La Histiocitosis de células de Langerhans (HCL) es una enfermedad de etiología y patogenia aún desconocidas. Afecta diferentes órganos y tejidos en los que produce lesiones de distinta gravedad. La histopatología de las lesiones y la clínica sugieren la participación de citoquinas en su patogenia. La IL-1β podría tener un rol importante en el desarrollo de la enfermedad.

El objetivo de este estudio fue determinar las concentraciones de IL-1β de las salivas de pacientes pediátricos con diagnóstico de Histiocitosis de Célula de Langerhans con y sin manifestaciones bucales (grupos 1 y 2 respectivamente), en relación a un grupo control (grupo 3), de pacientes pediátricos que no presentaron antecedentes médicos ni lesiones bucales.

Se estudiaron las salivas de 20 pacientes con la enfermedad de HCL, en relación a un grupo control de 11 pacientes pediátricos que no presentaron antecedentes. Los niños con Histiocitosis, con edades que oscilaban entre 4 meses y 16 años, fueron derivados del servicio de Oncohematología del Hospital Garrahan y Hospital de Clínicas, a la Cátedra de Odontología Integral Niños de la Facultad de Odontología de la Universidad de Buenos Aires. El análisis de los resultados se realizó según el test de Kruskall Wallis, se obtuvieron diferencias significativas entre los tres grupos (H = 20.36; P < 0.001). Luego se realizó el análisis de comparaciones múltiples de Dunn que mostró diferencias estadísticamente significativas entre los grupos 1 y 2, y entre los grupos 1 y 3 (p < 0.05). Se observaron valores más elevados de IL-1β en los pacientes con Histiocitosis con oral manifestations (Grupo 1) en relación con los pacientes sin manifestaciones (Grupo 2) y en el control del Grupo 3.

Palabras clave: saliva, Histiocitosis, Langerhans cells, interleukin, 1beta, mouth disease, child.

CONCENTRACION DE IL-1β EN SALIVA DE NIÑOS CON LESIONES BUALES ASOCIADAS A HISTIOCYTOSIS

RESUMEN
La Histiocitosis de células de Langerhans (HCL) es una enfermedad de etiología y patogenia aún desconocidas. Afecta diferentes órganos y tejidos en los que produce lesiones de distinta gravedad. La histopatología de las lesiones y la clínica sugieren la participación de citoquinas en su etiología. La IL-1β podría tener un rol importante en el desarrollo de la enfermedad. El objetivo de este estudio fue determinar las concentraciones de IL-1β en saliva de pacientes pediátricos con diagnóstico de Histiocitosis de Célula de Langerhans con y sin manifestaciones bucales (grupos 1 y 2 respectivamente), en relación a un grupo control (grupo 3), de pacientes pediátricos que no presentaron antecedentes médicos ni lesiones bucales. Se observaron valores más elevados de IL-1β en los pacientes con Histiocitosis con oral manifestations (Grupo 1) en relación con los pacientes sin manifestaciones (Grupo 2) y en el control del Grupo 3.

INTRODUCTION
Langerhans Cell Histiocytosis (LCH) is a disease whose etiology and pathogenesis are still unknown. It is characterized by a proliferation of histiocytes, lymphocytes and eosinophilic cells that form clusters producing several types of lesions. Different organs and tissues are affected, mainly skin, bones, lungs and liver. Bone tissue is affected...
in 72% of pediatric patients. The most compromised parts are cranium (27%), femur (13%), jaws (11%), pelvis (10%), vertebral bodies (8%), ribs (8%), humerus (5%), and tibia (3%). Its presentation may vary from a single bone lytic lesion with possible self-resolution without therapeutic intervention, to a disseminated form requiring medical treatment, corticoid administration, and/or chemotherapy.

In addition, lesions may appear within the oral cavity. The most frequent are early dental eruption, dental loss, gingivitis, periodontitis, gingival enlargements and gingival bleeding. The anatomical-pathological study of histiocytic lesions shows presence of Langerhans cells, which are gigantic multinucleated histiocytes similar to osteoclasts. Electron microscope images show intra-cytoplasmic inclusions: tennis racket shaped granules, called Birbeck granules, which are specific to pathological Langerhans cells. Their presence confirms diagnosis of LCH.

The diagnosis is completed through histochemical analysis to mark proteins CD1a and S 100 for these cells. The etiology and pathogenesis of this disease are still unknown; however, there are numerous hypotheses about its origin, including infectious, immunological, genetic, and oncologic. The immunological theory is one of the most convincing, because histologically the disease shows similarities to diseases of immune-reactive nature, and patients have diminished suppressing lymphocytes (CD8). In addition, the immunological alteration may be regulated by lymphokines and other growth factors that might modify the migration and maturation of Langerhans cells.

The pathogenesis of LCH is unknown and there is controversy regarding whether it is a reactive or a neoplastic process. LCH is not recognized as a malignancy. Despite the existence of cell cloning, the disease does not spread through the lymphatic system, and no metastasis occurs.

After an indeterminate period of time, reactivations (new isolated lesions) appear in most patients. The reactivations may or may not differ from the initial lesion, and may even involve different tissues. Despite the unknown origin of LCH, what is known is that many immunological mechanisms are involved in its pathogenesis, with the clinical picture and lesion histopathology suggesting that cytokines may participate in it.

### Classification
Lichtenstein (1853) proposed the term Histioctyosis X to group three clinical entities: Eosinophilic granuloma, Hand–Schüller–Christian’s disease, and Letterer–Süwe’s disease, adding the letter X to them because he did not know their etiology.

In 1973, Neselof et al. proposed Langerhans cells (LC) as responsible for the disease, which they called Langerhans Cell Histiocytosis (LCH). To systematize the disease according to the organs involved and further treatment, in 1985 the Histiocyte Society, an international scientific entity, proposed the present classification, as follows: Unifocal single-system histiocytosis: involves a single organ or tissue, usually bone or skin, and has a single location (focus). Corresponds to eosinophilic granuloma in the former classification. Multifocal single-system histiocytosis: involves a single organ or tissue, with multiple foci. Corresponds to Hand–Schüller–Christian’s disease in the former classification. Multisystem histiocytosis: involves multiple organs or tissues, with multiple foci. It is acute and usually occurs in infants from birth. Corresponds to Letterer–Süwe’s disease in the former classification.

The aim of this study was to determine the concentrations of IL-1β in saliva of pediatric patients diagnosed with Langerhans Cell Histiocytosis, with and without associated oral histiocytosis lesions.

### MATERIALS AND METHODS
The sample consisted of a total 31 pediatric patients, who were divided into three groups:
- **Group 1**: 10 patients with Langerhans Cell Histiocytosis, with lesions in the oral cavity.
- **Group 2**: 10 patients with Langerhans Cell Histiocytosis without oral lesions.
- **Group 3**: 11 patients with neither medical antecedents nor oral lesions.

Patients in Groups 1 and 2 were referred by the Oncohaematologic Service, Garrahan Hospital, diagnosed with Langerhans Cell Histiocytosis. Group 3 consisted of healthy pediatric patients who visited the Department of Comprehensive Children’s Dentistry, School of Dentistry, University of Buenos Aires (UBA), for dental assistance. The inclusion criteria for Groups 1 and 2 were confirmed diagnosis of Histiocytosis, having signs –
symptoms compatible with LCH seen by light microscopy, plus confirmation by electron microscopy of presence of Birbeck’s granules and positive immuno-tracing with CD1a in the lesion cells (Table 1).

Patients taking any medication were excluded from all three Groups (Table 1).

**Methodology**

1. Parent’s informed consent for performing this study was obtained. The study was accepted by The Ethics Committee of the School of Dentistry.
2. Dental files and history of the disease in each child were made.
3. Lesions of the oral mucosa were located and diagnosed through visual examination and palpation. Biopsies were sent to Department of Pathological Anatomy, School of Dentistry, Buenos Aires University.
4. Periapical and panoramic radiographs were taken of each child in order to detect osteolytic lesions.
5. Samples of non-stimulated saliva from the three groups were collected in sterile tubes, covered, and sent to the Department of Pharmacology at the same School in order to establish the concentration of IL-1β. The samples were stored in a freezer at -80º until they were analyzed.

All the procedures used are specified in the protocol for diagnosis of IL-1β.

**RESULTS**

Values for Group 1 (with lesions in the mouth) ranged from 2,625.26 to 749 pg/ml, most of them higher than 1,000 pg/ml.

Values for Group 2 (without oral lesions) ranged from 53.84 to 503.44 pg/ml.

Values for Group 3 (control) ranged from 10 to 496.78 pg/ml.

Medians were 1,309.11 for Group 1; 243.54 for Group 2, and 139.00 for Group 3 (Control) (Table 2).
The Kruskall Wallis test applied to the results showed significant differences among the three groups ($H = 20.36; P < 0.001$).

Dunn’s multiple comparison analysis showed statistically significant differences between Groups 1 and 2, and between Groups 1 and 3 ($P < 0.05$) (Fig. 1). Values of IL-1β were significantly lower in patients presenting lesions outside the mouth, and with no significant difference compared to Group 3 (Control).

**DISCUSSION**

For over 40 years, saliva has been considered an auxiliary tool in the diagnosis of oral diseases, because organic molecules of proteinic nature have been detected in its composition, mainly specific antigens, proteinic cell particles – receptors, glycoproteins, cytokines, (interleukines and derivatives), which may be related to oral lesions. Serum components can also reach saliva through the gingival crevicular fluid, providing potential application in the diagnosis of certain disorders. The etiology of Langerhans Cell Histiocytosis has not yet been established; nevertheless, it is accepted that it is the expression of an immunological disorder.

Similarly, the pathogenesis of LCH is enigmatic, although the altered expression of cytokines and cellular adhesion molecules, which are important for migration and homing of the normal Langerhans cells (LC), may play an important role.

Kannourakis et al. (1994) and Egeler (1999) suggested that cytokines participate in the histopathology of the lesions. It has been observed that the production of cytokines plays an essential role in reactivations of the disease, as well as in the inflammatory and immunological processes. Specifically, the IL-1β are important in the development and evolution of Langerhans cells; and the Langerhans cells also produce them. Interleukines act on osteoclasts linked to bone resorption. IL – 1 was found in bone granulomas diagnosed as histiocytosis.

However, the regulation of cytokine production is still unknown. In other oral diseases, high cytokine values were found in saliva. Kaushik et al. (2011) found high levels of IL-1β in saliva of patients with untreated chronic periodontal disease (comparable to those found in this study in children with oral lesions), and those levels decreased significantly after basic periodontal therapy. Katakura et al. (2007) reported a significant increase in cytokines IL-1β and IL-16 in serum and saliva of patients with oral cancer compared to the healthy Control Group. However, no previous study analyzing saliva of patients with histiocytosis was found.

In the present study, among the patients with Langerhans Cell Histiocytosis who showed oral lesions, eight were primary and two were reactivations. Among the oral manifestations, Hernandez and Juyol et al., found mandibular osteolysis, which can lead to a reduction in mandible height and bone loss linked to inflamed gingiva, looking like “floating teeth”. The “floating tooth” is a pathognomonic sign of the disease.

In this study, the lesions in mucosa in Group 1 clinically look like erosions (Fig. 2), and osteolytic lesions of support bone associated to atypical enlargements of oral mucosa were observed (Figs. 3 and 4). Solitary osteolytic lesions in maxillary bone (Fig. 5), bone reactivations with localized bone loss, similar to periodontal disease (Fig. 6), and bone lesions located in deciduous teeth (“floating tooth”) were observed (Fig. 7).

In this study, open lesions were diagnosed in the oral cavity with inflammatory exudates and closed jaw injuries without any exudation. In all cases, high IL-1β values were found in saliva.
Figs. 2 to 7 are clinical examples showing the diversity and aggressiveness of oral lesions associated with LCH.

The panoramic radiograph (orthopantomograph) is the method of choice to study the jaws. It allows observation of the integrity of the cortex, presence of osteolytic lesions in maxilla and mandible that may compromise tooth buds, and comparisons with later radiographs during the follow-up of each patient9,10. The active bony lesions present indefinite borders, and are rounded or elliptic. In contrast, when the lesion is healing, the size and density of the trabecular bone within the lytic areas diminishes, and the thickness of the lesion margin increases, appearing as a radiopaque halo similar to cortex bone43. The high levels of IL-1β found in saliva of children with histiocytosis lesions in the oral cavity
allow us to assess or raise the value of saliva as a fluid that could contribute to the diagnosis of LCH. It is concluded that:

- Pediatric patients with Langerhans Cell Histiocytosis with oral manifestations show high values of IL-1β in saliva, which are higher than those in patients who suffered the disease without oral manifestation, and higher than those in the group of healthy children.

- A saliva sample, which is a simple, non-invasive procedure, may allow detection of high concentrations of IL-1β associated to primary or reactivation lesions in the mouths of children with Langerhans Cell Histiocytosis.

- Further studies should be conducted on a larger sample to confirm the association between the oral lesions and the interleukines in saliva from patients with LCH.

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REFERENCES


