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Case Report

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Diagnosis Procedure In Case Of a Gingival Hyperplasia: Presentation of One Case

Kwedi K.G.G^{1.2}, Diakite Y^{1.2}, Kane A.S.T², Fokwa N.G¹, Daramsis H^{1.2}, Messina E.W¹, Diatta M³, Tamba B³, Dia Tine S³

¹ Specialized degree of Oral surgery, Institute of Odontology and Stomatology, University Cheikh Anta Diop, Dakar, Senegal

² Master in Odontological Sciences, Institute of Odontology and Stomatology, University Cheikh Anta Diop, Dakar, Senegal

³ Oral Surgery Department, Institute of Odontology and Stomatology, University Cheikh Anta Diop, Dakar, Senegal

*Corresponding Author

Kwedi K.G.G

Abstract: Introduction: Gingival hyperplasias are abnormal increases in gingival volume, due to exaggerated proliferation of its cells. There are a number of causes for this condition. The cause can be local but also be the expression of certain hormonal status, of general pathologies and/or their therapies. **Observation:** A 12 year old patient, suffering from anemia, consulted for a painful gingival overgrowth evolving for more than a year. The diagnosis procedure led us to an inflammatory gingival hyperplasia of infectious origin. **Discussion:** In the presence of gingival hyperplasias the case history and the clinical examination orientate the diagnosis. Further examinations permit to retain the cause and exclude other possible etiologies. This procedure can turn out difficult. **Conclusion:** Before the multiple etiologies possible for a gingival hyperplasia in a child, it is important to carry out complematary examinations in order to avoid bad surprises.

Keywords: Diagnostic approach, gingival hyperplasia, clinical case.

INTRODUCTION

Gingival hyperplasia is an overgrowth of gum tissue as a result of abnormal (excessive) proliferation of its cells. There are a number of causes for this condition. They can be inflammation causes, hormonal (puberty, pregnancy, hypothyrodism), systematic causes (vitamin deficiency), tumorous (benign tumour of the buccal mucosa, proliferative hemopathia) or be druginduced as side effect of prescribed medications (calcium inhibitors, anticonvulsants, immunosuppressors) (Persson, R.E. *et al.*, 2003; Dongari-baqtzoglou, A. 2004; Laine, M.A. 2002).

In case of gingival hyperplasia in a child, the case history and the clinical examination give different diagnosis orientations. Further examinations permit, after exclusion of all other etiologies, to retain a diagnosis.

We present here by, the diagnosis procedure in case of a gingival hyperplasia.

A 12 year old patient addressed to the stomatology service of the general hospital of Grand Yoff at Dakar-Sénégal by his treating dental surgeon for painful gingival over swelling located at the mandibular at the mandibular. According to the patient, the lesion has been evolving for more than a year and had incressed in size with time. The patient described major pain on teeth contact and/or during feeding, associated with weight loss. Six months earlier, she consulted her treating medical doctor for asthenia and anorexia. The biological examination revealed a severe anaemia, a blood transfusion was done after patient hospitalisation. No past medical history was retained.

Clinical Observation

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Figure 1: Right mandibular gingival hyperplasia.

At the exobuccal examination, we noted the presence of soft, submandibular bilateral adenopathies, mobiles, unpainful on palpation. Endobuccal clinical examination revealed the presence of a gingival overswelling going from the 31 to 47. The lesion with erythematous aspect occupied the vestibular and lingual borders of the mentioned teeth with partial colonization of the occlusal surfaces. The lesion was soft, bleeding on contact (fig. 1). The mucosa covering the buccal cavity was pale colored. The presence of tartar was noticed at the first and fourth quadrant, testifying of a preferential left unilateral mastication. At the end of the first consultation, several diagnosis hypotheses were formulated: inflmatory gingival hyperplasia, epulis, proliferative hemopathia. A radiological examination (orthopantomogram) and biological examinations (complete blood count, coagulation factors) were requested.

Table I : Results of blood count. Values followed bya * are anormal.

	Results	Unit	References
Erythrocytes	4,00	T/L	4,2 - 5,2
Haemoglobine	11,4*	G/dL	12 - 16
Plaquettes	497*	G/L	150 - 450
Procalcitonin	0,50*	mg/L	0,1 - 0,4
PN neutrophiles	2,63	G/L	4,2 - 5,2
PN basophiles	0,14*	G/L	< 0,10
Lymphocytes	2,35	G/L	1 - 4
Monocytes	0,71	G/L	0,2 – 1
VS	40*	mm.1h	<20



Figure 2: Initial Orthopantomogram

The orthopantomogram revealed a discret print of the mower dental canal, a mesioversion of the 45 with a slight bone lysis between 45and 46 (fig. 2). The complete blood count put in evidence an anaemia (11, 4 g/dl), a thrombocytosis (497 giga/L), a basophilia (0, 14 giga/L) and an increase in procalcitonine (0, 50%). The sédimentation speed was of 40 mm within him first hour (Tab. I).most of these tests was in favor of an infectious process. However the recurrent anaemia and the basophilia made us fear an atypical proliferative hemopathia. Before this clinical picture, a biopsy was realized during this consultation, this with objective the exclusion hypothesis of an atypical proliferative hemopathia. The histological examination of the biopsy revealed Malpighian epithelia presenting a slight acanthosis, keratosic without cytopathy nor atypical cytonuclear.



Figure 3: Healing of the gingiva six months later.

The clinical and complementary examinations permitted an exclusion of other hypothesis. The only plausible diagnosis amongst the ones evoked was that of inflammatory gingival hyperplasia of infectious origin. A medical prescription (amoxicilline + clavulanic acid 2g, 2x/day; paracetamol 3g, 3x/day) and a mouthwash made of Chlorhexidine were given. A scaling and polishing followed of a resection of the hyperplasia with a cold bistoury blade; then a cautérisation with the electrical bistoury under local anaesthesia were realised on the dental chair. Control was done at day 7, day 14, day 30, day 60, day 120 and day180 (**Fig. 3**). The patient was sent back to the haematolgy service for specified examinations and the handling of her anaemia.

DISCUSSION

We report here a case of inflammatory gingival hyperplasia lasting over a year on a young girl suffering from anaemia with no past history of medication known.

Clinically, inflammatory gingival hyperplasia presents an oedematous, shinning red purplish soft gingival that easily bleeds. The voluminous gingival mostly covers and important part of the crowns, leading to an increase in the sulcus depth, favoring sticking of food debris and accumulation of bacteria (kinane, D.F. 2001). Inflammatory gingival hyperplasia is mostly of local cause, provoked by bacterial toxins inducing biofilm through its enzymes and toxines induising an inflammatory response of the gingiva. Nevertheless, certain cases of inflammatory gingival hyperplasia due to buccal respiration have been reported in literature (Laine, M.A. 2002; kinane, D.F. 2001).

Histologically, gingival hyperplasia is composed of, an excess of fibrous conjunctive tissue covered by a squamous stratified epithelium of normal thickness or hyperacanthosic (thickening of the mucous body of malpighi) (Kaqueler JC et Le May O 1998).

The positive diagnosis must be done, excluding all other causes of gingival hyperplasia. In our case, epulis and proliferative hemopathia were cited as differentials.

Gingival epulis is amongst the gingival hyperplasic pseudo-tumours. The most frequent causes cited are traumatic and hormonal. We find them at the level of the neck of one or two contiguous teeth. It can be sessile pediculated or ligamented, staying classically on the incisivo-canin vestibular section (Ch. FATY N'DIAYE, B. *et al.*, 1996; AKAZANE, A., & BADREDINE, H. (2014). In our case, gingival hyperplasia occupied the vestibular and lingual sectors going from 31 to 47.

Proliferative hemopathia are the most frequent malignant pathologies found in children less than 15 years old (Sepúlveda, E. *et al.*, 2012). In certain aspects, this case ressembled a proliferative hemopathia:

- The first, frequently seen gingival hyperplasia that usually results from the infiltration of oral tissues by malignant cells;
- > The second is the presence of anaemia ;
- The third corresponds to the increase in the amount of polynuclear basophiles circulatingwhich is observed in cases of proliferative hemopathia, chronic haemolytic anaemia (Sepúlveda, E. *et al.*, 2012; Rerhrhaye, M. *et al.*, 2010).

However, there was no uneplained dental pain at the clinical exam and the blood count returned without neutropenia, nor thrombopenia. Elements frequently seen in cases of proliferative hemopathia (Sepúlveda, E. *et al.*, 2012; Zhou, Y. *et al.*, 2011). Several stages of evolution and of gravity exist, from the most discret to the most evident. Moreover, certain cases of atypia have been described in literature, making the diagnosis more difficult (Lafon, A. *et al.*, 2010). The coagulation factors came without any particular informations. We noted an increase in the procalcitonin concentration. Procalcitonin is a prohormone which blood concentration can be measured specifically. Its increase in blood is revelatory of an inflammatory process, precisely bacterial, this, even in the absence of neutropenia (Sager, R. *et al.*, 2017).

Gingival hyperplasia is less frequent in children. Granulomatous diseases (Crohn'sdisease, Wegener granulomatosis), malignous tumours and hereditary must be taken into consideration in the differential diagnosis of gingival hyperplasia in children (Krishna, K. B. *et al.*, 2014; Olczak-Kowalczyk, D. *et al.*, 2011). In our case, the diagnosis retained after a synthesis of all collected elements was that of inflammatory gingival hyperplasia of infectious origin.

When the hyperplasia is minimal, a careful scaling of the teeth associated with a correct hygiene can be sufficient. For the important lesions, a rigorous surgical intervention is necessary (Cuest, A. *et al.*, 1998). In our case, the treatment associated a cleaning of the buccal cavity under antibiotic covering and a gingival resection under local anaesthesia.

Recidivism is the most common problem in the management of gingival hyperplasia. To avoid this recidivisms, it is recommended to program a gingival therapy of maintenance (Cuest, A. *et al.*, 1998). So we reviewed our patient for control sessions. Six months later the patient was doing great and the gingival mucosal looked stable.

CONCLUSION

The objective of this work was to demonstrate that for a gingival hyperplasia in a child we can have different diagnosis. Before the multiple etiologies possible gingival in a child, certain being able to involve the vital pronostic of the patient, it is important to do complematary examinations in order to avoid bad surprises.

Conflict of Interest

None.

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