**Abstract**

Orofacial granulomatosis is a chronic granulomatous condition characterized by relapsing and remitting lip swelling and oral involvement that may include deep ulcers, tags and cobblestone formation. It occurs as an independent entity but also in conjunction with systemic diseases such as tuberculosis, sarcoidosis and Crohn’s disease. The clinical presentation is not indicative of concomitant systemic disease. To highlight the importance of thorough examination to rule out systemic disease, we present two childhood cases of orofacial granulomatosis, one of which was associated to Crohn’s disease.

**Case Report #1**

Case 1 was a 12-year old boy referred to us because of persistent swelling of the lips and gingivae, along with an uncharacteristic facial rash (Figure 1). He had a history of atopic dermatitis, but was otherwise previously healthy and had no predisposition to skin- or gastrointestinal disease. He presented with pronounced perioral edema, severe fissuring of the lips, angular cheilitis, gingival edema, and cobblestone formation of the buccal mucosa. During flare-ups the boy experienced severe abdominal pain. Suspecting inflammatory bowel disease, the patient was thoroughly examined with blood tests, biopsies from the gingival and buccal mucosa, and endoscopy of the gastrointestinal tract. The histopathological examination of the mucosal biopsies revealed chronic granulomatous inflammation consistent with OFG and oral Crohn’s disease. Serum calprotectin was elevated to 409 mg/kg. CRP and blood sedimentation rate were normal. Gastroscopy showed duodenal inflammation and by subsequent endoscopy ileač aphthose ulcers were detected. The patient was diagnosed with Crohn’s disease. Infliximab treatment was initiated with a good clinical response on both abdominal symptoms and mucosal lesions.

**Case Report #2**

Case 2 was a 9-year old boy who presented chronic swelling of the upper lip, redness, swelling and hypertrophy of the gingivae, and cobblestone formation of the buccal mucosa (Figure 2). This patient had IgE-mediated allergy to grass, birch, and alternaria, but was otherwise healthy and was not predisposed to any skin or systemic diseases. He had no history of abdominal pain or stool alteration and no respiratory symptoms. As in case 1, the mucosal biopsies from the upper-lip revealed granulomatous inflammation. All blood tests were normal. The patient was further conferred with a pediatric gastroenterologist. Serum and fecal calprotectin were normal. As this patient had no history of abdominal problems, no further examination was indicated at the time. The oral lesions were treated with topical fluticasone propionate 50 g/dose. The initial dosage was 1 spray unit twice daily. After 6 weeks this was reduced to once daily for another 6 weeks, after which the treatment was given pro necessitate. The lesions responded well to this treatment and after 12 weeks only mild infiltration and swelling of the gingivae remained without noticeable inflammatory activity. Regular check-ups every 3 months ensure that the condition is stable, and that possible gastrointestinal symptoms are detected.

**Discussion**

Whether OFG should be considered a separate entity or a manifestation of systemic disease remains a topic of discussion. In particular, several case reports have suggested an association to Crohn’s disease. The two entities share a number of clinical and histological features, but the exact relationship has not yet been established. Some authors suggest that patients with OFG may have subclinical Crohn’s disease. Furthermore, several authors have observed that children are more likely to have onset of OFG preceding symptoms of Crohn’s disease, and speculate that childhood onset carries a higher risk of developing Crohn’s disease.

The prevalence of Crohn’s disease in children with OFG is not well-established and could be underestimated since our current knowledge is based on case reports and case studies. The clinical presentation of the two patients presented here was almost identical. The history revealed that one of the patients had intermittent abdominal pain, which led to further examination, revealing an underlying Crohn’s disease. This illustrates an important point, to be remembered when encountering children with OFG. The clinical presentation is not indicative of concomitant systemic disease and symptoms of this may be few. Regardless of the clinical presentation, systemic disease should always be ruled out. OFG has been reported to be a manifestation of tuberculosis and sarcoidosis however, the vast majority of the literature focuses on the possible link...
Case Report

Table 1. Differential diagnoses of orofacial granulomatosis.

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>Granulomatous swelling of lips</th>
<th>Cobblestone formation of oral mucosa</th>
<th>Mucosal swelling</th>
<th>Oral ulcers</th>
<th>Facial nerve paralysis</th>
<th>Plicated tongue</th>
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</thead>
<tbody>
<tr>
<td>Orofacial granulomatosis</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>(+)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Granulomatous cheilitis</td>
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<td>-</td>
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<tr>
<td>Melkersson-Rosenthal Syndrome</td>
<td>+</td>
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between OFG and crohn’s disease (Supplementary Table 1)."3,11,12,14,16 In our opinion, the caring for pediatric patients with OFG should focus mainly on monitoring for signs, symptoms, and laboratory evidence of crohn’s disease. A thorough patient history and clinical examination is necessary and we recommend blood screening including CRP, blood sedimentation rate, complete blood count, and serum calprotectin. If serum calprotectin is elevated, the patient should be referred to a pediatric gastroenterologist. Any specific symptoms or abnormal blood screening should lead to further examination of relevance, i.e. a history of pulmonary symptoms should lead to X-ray examination of the chest and further blood tests depending on the symptoms and patient history. We recommend that patients be followed closely until the oral symptoms stabilize. When the condition is stable follow up at regular intervals ensures that the condition remains stable and any new symptom of systemic disease is detected.

References