The Silent Sinus Syndrome: A Collaborative Approach between Rhinologists and Oculoplastics- Case Report and Literature Review

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Abstract

The Silent Sinus Syndrome (SSS) is a rare phenomenon originally described in 1964 by Montgomery, which often occurs unilaterally in the maxillary sinus with opacification and collapse. It is characterized by ipsilateral enophthalmos and hypoglobus or mistakenly as an exophthalmos of the contralateral eye. The pathophysiology is caused by obstruction of the ostiomeatal complex with subsequent maxillary sinus hypoventilation and development of negative intra-sinus pressure resulting in bony changes and sinus collapse. Radiographic evaluation by the mean of CT scan orbits and paranasal sinuses as well as nasal endoscopic examination are mandatory in diagnosing such cases. Treating such cases requires a collaborative multidisciplinary approach to ensure the best possible results. In this case report, we discuss this rare phenomenon in terms of clinical presentation, clinical and radiological evaluation and conclude with the optimal treatment approach.

Keywords: Silent sinus syndrome; Enophthalmos; Hypoglobus; Ostiomeatal complex; Maxillary sinus

Introduction

Silent Sinus Syndrome (SSS) is a rare clinical disorder that is typically characterized as spontaneous, painless, progressive enophthalmos and hypoglobus. It results from downward bowing of the orbital floor secondary to maxillary sinus collapse, in the absence of any symptoms of sinonasal disease [1]. In 1964, Montgomery was the first to present and publish two cases of unilateral enophthalmos and hypoglobus associated with ipsilateral maxillary sinus collapse; however, SSS term was only used 30 years later by Soparkar and colleagues [1,2]. Interestingly, the term imploding antrum is another term used for silent sinus syndrome in literatures review [3]. Recent to this in 2008, Brandt and wright suggested SSS to be included under the general term of chronic maxillary atelectasis [4]. It has been noticed that increasing numbers of case reports were published on SSS recently, which could be explained by the widespread use of Computed Tomography (CT) and endoscopic interventions in diagnosing such cases. This case report will illustrate the clinical and the imaging features of this rare phenomenon. Our aim from this report is to emphasize in the collaborative approach required to manage such patients with this syndrome. Additionally, this report will highlight the significance of rare but not to be missed differential diagnosis in any unexplained enophthalmos, hypoglobus or even exphthalmas. On further questioning the patient, he denied having pain, loss of vision, rhinorrea or any history of facial trauma. Moreover, he denied any past ophthalmic or sino-nasal surgeries. Careful ophthalmic examination revealed his Best -Corrected Visual Acuity (BCVA) to be 20/25 of the Right Eye (RE) and 20/30 of the Left Eye (LE). Exophthalmometer measurement of the RE was 17 mm compared to 14 mm for the LE (3 mm difference) and facial asymmetry was noticed. Full ocular movements were intact with normal findings on anterior and posterior segments’ examination. Initial diagnosis was RE exophthalmos; hence, thyroid function test was ordered with 3 weeks follow-up appointment given. The patient was seen in the follow-up visit but this time with new symptoms and complaints (Figure 1) as follow:

- More diplopia in primary gaze.
- Worse proptosis (4 mm) of the RE
- More pronounced facial asymmetry.
- Deep upper lid sulcus of the LE
- LE hypoglobus.

Case Report

A 47-year-old, previously healthy, Caucasian presented to the orbital and oculoplastics clinic complaining of 2 weeks duration of occasional binocular vertical diplopia associated with right eye exophthalmos. On further questioning the patient, he denied having pain, loss of vision, rhinorrea or any history of facial trauma. Moreover, he denied any past ophthalmic or sino-nasal surgeries. Careful ophthalmic examination revealed his Best -Corrected Visual Acuity (BCVA) to be 20/25 of the Right Eye (RE) and 20/30 of the Left Eye (LE). Exophthalmometer measurement of the RE was 17 mm compared to 14 mm for the LE (3 mm difference) and facial asymmetry was noticed. Full ocular movements were intact with normal findings on anterior and posterior segments’ examination. Initial diagnosis was RE exophthalmos; hence, thyroid function test was ordered with 3 weeks follow-up appointment given. The patient was seen in the follow-up visit but this time with new symptoms and complaints (Figure 1) as follow:

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Consequently, axial and coronal CT scan images of the orbits and paranasal sinuses (Figures 2 and 3) were ordered. Radiological findings
showed partial opacification of the left maxillary sinus with reduced maxillary sinus volume and ipsilateral globe prolapse associated with almost complete bony resorption of the left orbital floor. All of these findings were consistent with the diagnosis of Silent Sinus Syndrome.

To ensure the best possible outcomes, the decision was made to do Combined Functional Endoscopic Sinus Surgery (FESS) with orbital floor reconstruction under the same anaesthesia and the same surgical table in a teamwork approach. FESS, in the form of uncinectomy and middle meatal antrostomy was performed. The left maxillary sinus was filled with secretions and polypoidal mucosa. Following the rhinologist’s intervention, oculoplastic took over performing anterior orbitotomy via transconjuctival approach with lateral canthotomy-cantholysis. Due to significant orbital floor loss, a Synpor titanium mesh 43 mm (radius) X 15 mm was fashioned and implanted to support the globe inferiorly. The histopathological examination showed non-specific chronic inflammation within the sinus mucosa. Bacterial and fungal cultures were negative. The patient was followed up for 12 months and showed dramatic improvement of the initially reported symptoms and signs. The patient reported no diplopia and thorough ophthalmic examination revealed good restoration of the upper lid sulcus with no signs of enophthalmos or hypoglobus (Figures 4 and 5) show almost normal maxillary sinus volume with restored orbital architecture supported by the titanium mesh implant in the left orbital floor.

Discussion

Upon reviewing literatures, several case reports have explained different presentations of this phenomenon; however, they all shared the combination of enophthalmos and hypoglobus in the presenting complaint. Due to its insidious course of presentation with no linked past history; SSS poses diagnostic challenge to otolaryngologists, ophthalmologists as well as general practitioners. When Soparkar et al. published his observations of SSS; only 30% of the cases had past history of sinus disease in childhood [5]. SSS is found to affect adults between the third and the fifth decades of life with no gender variation [4]. Additionally, experts of this phenomenon have found no association between smoking, alcohol consumption and genetics with the development of this rare syndrome [6]. The most common reported complaint to an ophthalmologist is changes to the facial appearance experienced by the patient or noticed by friends or family members [7]. It presents typically as unilateral asymmetry and can be sometimes misinterpreted as exophthalmos of the contralateral eye. According to Stevens et al. [3], approximately 50% of patients with enophthalmos are initially referred to ophthalmologists for investigations of contralateral exophthalmos or ptosis. Other presenting symptoms include lid lag, lagophthalmos, oscillopsia and transient vertical diplopia [8]. While enophthalmos is present in all cases, diplopia occurs in about one - quarter and it is typically vertical
and painless [7,9]. These orbital symptoms can suggest a possible neuro-ophtalmic disorder; however, the presence of enophthalmos with hypoglobus in previously healthy patients will narrow the differential diagnosis towards Silent sinus syndrome. In order to diagnose SSS; other causes that might present with similar symptoms must be ruled out first such as Horner’s syndrome, facial hemiatrophy, developmental anomalies or any past history of facial trauma [10]. In our case, the history was negative for the mentioned above disorders and his endocrine and autoimmune markers were negative too. Many theories have explained the pathophysiological process of this disease with varying degrees of evidence in literature. Gillman et al. in 1999 described the most acceptable theory behind the disease process [11]. Gillman et al. suggested that in SSS patients, obstruction of the Osteo Mental Complex (OMC) leads into hypoventilation of the maxillary sinus with the subsequent development of intra-sinus negative pressure and thinning of the sinus wall. This chronically causes remodeling and bone resorption of the orbital floor causing malposition of the ipsilateral eye. Although this theory is widely accepted among observers of this phenomenon; however, no one yet knows the reason behind OMC obstruction [4]. Radiographic evaluation by the mean of CT scan orbits and paranasal sinuses is crucial to assess the anatomy of orbito-maxillary sinus structure and is mandatory upon labeling patients with such a disease.

Table 1: Shows most clinical manifestations of SSS.

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>Value</th>
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<tbody>
<tr>
<td>Enophthalmos</td>
<td>ranging from 1-6mm</td>
</tr>
<tr>
<td>Hypoglobus</td>
<td>ranging from 0-6mm</td>
</tr>
<tr>
<td>Lid retraction</td>
<td>mistakenly diagnosed</td>
</tr>
<tr>
<td>Deeping of superior sulcus Exophthalmos</td>
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CT scan findings show opacification (complete or partial) of the maxillary sinus and collapse with inferior bowing of the orbital floor towards affected maxillary sinus [12]. Cobb et al. further explains CT scan findings by linking the pathophysiological process to the radiological findings, ‘the maxillary sinus roof loses its normal upward convexity and the orbital floor is bowed downwards giving the appearance of reduced sinus volume with increased orbital volume’. This orbital volume expansion causes horizontal displacement of the eye level appearing as hypoglobus. Treating such cases require a collaborative multidisciplinary approach to ensure the best possible results. It involves the input of rhinology, oculoplastics as well as anesthesia with the availability of good nursing team in the operating theatre. Treatment from the otolaryngology points of view is aimed at restoring drainage and maintaining good aeration of the occluded maxillary sinus. Traditionally, the Caldwell-Luc procedure used to be the gold standard in treating orbital floor. The use of titanium mesh fixed with Leibinger screws is the preferred material in reconstructing and maintaining the normal orbital architecture, classically through transconjunctival or external lower lid approaches [4,15]. Again, it is important to emphasis that the decision to perform both combined procedures should take the teamwork approach with the patient’s view took in great consideration while weighing up the risks and benefits of such a decision.

Conclusion
Silent sinus syndrome is rare and often misdiagnosed pathology with a unilateral progressive and asymptomatic collapse of the maxillary sinus. The pathogenesis is not completely understood but appears to result from maxillary sinus collapse due to acquired obstruction of the maxillary sinus drainage. CT scan is usually required for a definite diagnosis. Treatment of such a disease consists of restoration of maxillary sinus aeration and reconstruction of the orbital floor in case needed. Any unexplained enophthalmos especially if associated with hypoglobus should hint for silent sinus syndrome as a differential diagnosis.

References
