Epiphora as a sign of unexpected underlying squamous cell carcinoma within sinonasal inverted papilloma

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Abstract.

Background: Sinonasal inverted papilloma (SIP) is a benign tumor of the nasal cavity and sinus. It is characterized by aggressive malignant transformation and a high rate of recurrence. Inadequate removal of the tumor during surgery is one of the most significant contributors to recurrence. SIP has been reported to rarely occur in conjunction with Squamous Cell Carcinoma (SCC) invading the lacrimal drainage system and the orbit. The mechanism of this secondary SCC transformation has yet to be explained.

Case presentation: Herein, the authors present the case of a 66-year-old woman with a rapidly enlarging sinonasal inverted papilloma with secondary squamous cell carcinoma of the right nasal cavity presenting with epiphora.

Conclusions: Nasolacrimal duct obstructions require a thorough examination as it may hide a malignant pathology.

Keywords: sinonasal inverted papilloma; squamous cell carcinoma; oculoplastic surgery

Introduction.

SINonasal inverted papilloma (SIP) is a benign tumor which originates from the sinonasal Schneiderian mucosa and accounts for 0.5 to 4% of all nasal and sinus neoplasms. [1] Pathologically, SIP epithelium inverts into submucosal stroma, which is different from other types of nasal papilloma. Unlike other benign tumors, SIP exhibits remarkable aggressive behaviors, including invasiveness, recurrence and malignant transformation. [2] Therefore, SIP can spread into the paranasal sinus, orbit, and cranial base, which can lead to poor prognosis for SIP patients. [2] Secondary squamous cell carcinoma is malignancy formation within the SIP. It is considered primary if there is no additional mucosal condition or secondary if it occurs in conjunction with an additional condition, which is estimated to occur in about 9% of cases. [3] The authors report the case of a 66-year-old woman with SIP and secondary SCC of the right nasal cavity. The unique feature is the scarce symptoms related to the advanced disease stage. This study was adherent with the tenets of the Declaration of Helsinki.

Case presentation.

A 66-year-old woman presented with a 4-year history of right-sided epiphora and new onset swelling at the right medial canthus mimicking a lacrimal sac’s mucocele. She denied bloody tears, pain and diplopia. The patient was referred to the hospital eye service with a diagnosis of a chronic dacryocystitis and nasal obstruction. The ophthalmologic examination revealed a firm, non-tender, 1 cm mass palpable just below the right medial canthal ligament. Irrigation showed regurgitation of mucopurulent material. Uncorrected visual acuity was 7/9 in both eyes. No RAPD nor proptosis were detected. No other palpable masses nor lymphadenopathy were present in the head and neck region. Endoscopic examination of the nasal cavities showed a polyloid lesion of the right lateral wall and diffuse nonspecific mucosal edema. CT scan of the orbit revealed 2 x 2 cm well defined, heterogeneously enhancing abnormal soft tissue mass in the right nasolacrimal sac, duct and lateral wall of right nasal cavity. Bony erosion of the orbital rim was observed (Fig 1 A-B).
Figure 1A: Non enhanced axial CT shows bone erosion of the papyracea lamina of ethmoid bone.  
Figure 1B: Axial T2 weighted sequence shows an extraconal and intracanal mass (white arrows) in the right orbit, with non-homogeneous hyperintensity ed irregular margins. The lesion involves the nasolacrimal canal, the papyracea lamina and the anterior ethmoid cells.

MRI scan substantiated a T2 mildly hypointense, T1 isointense mass with erosion of the inferior orbital wall. Endoscopic biopsy was carried out and the histological examination detected an inverted papilloma with synchronous squamous cell carcinoma (Fig 2).

Fig. 2 - HE 10X and HE 40X  
Malignant squamous cell proliferation characterized by scanty keratinization, abundant and multifocal necrosis and sparse mitotic figures, organized in papillary and polypoid structures, partially cystic, closed to fibrous and osseous tissue.

A multidisciplinary approach is our preferred method of choice for cases with both nasal and orbital involvement. The surgical resection planning was a joint effort of the Ophthalmology and the Otorhinolaryngology departments.

The patient underwent lateral rhinotomy with orbitotomy. The orbital cavity was filled with a penduculated temporal muscle flap, covered with a dermo-epidermal skin graft (Thiersch graft). Histological examination reported radical tumor excision and immunohistochemical staining was negative for Ki67 and HPV.

The patient was submitted to adjuvant radiotherapy after surgery. The total dosage of radiation was 60 Gy in 30 sessions. Endoscopic examinations and medications of the nasal fossa were performed every three months in the follow-up period. One year after surgery, both PET and MRI of the head and neck were negative for recurrent disease. During the following 4 years of follow-up, MRI was performed together with clinical examinations and neck ultrasound. No recurrence was detected.

Discussion

Sinonasal Inverted papilloma (SIP) is a rare benign sinonasal tumor, three characteristics make this tumor very different from other sinonasal tumors: a relatively strong potential for local destruction, high rate of recurrence, and a risk of carcinomatous evolution. [4] Unlike most papillomas of the body, this mucosal lesion inverts, rather than everts, into the underlying connective tissue stroma. [1]
Clinical presentation is most commonly a male subject who is 50–69 years old. Classically, the “tumor” originates from the lateral nasal wall, particularly in the osteomeatal complex area of the middle meatus, and adjacent inferior and middle turbinate, and subsequently involves the contiguous paranasal sinuses, with the maxillary sinus as the most frequent sinus involved and ethmoid sinus as the second most common. Unilateral nasal obstruction and intermittent epistaxis are the most common presenting symptoms. [1]

Up to 90% of tumors of head and neck are squamous cell carcinomas (SCCs) [5], which represent the seventh most prevalent cancer worldwide. The male-to-female incidence ratio is 3:1, and about 70% of new cases occur in low- and middle-income countries. In 2018, there were an estimated 453,000 deaths from head and neck cancer globally. In the past 15 years, strong evidence has accumulated that infection with certain human papillomaviruses (HPVs) is etiologically involved in a subset of head and neck cancers. [6]

Squamous cell carcinoma may be a secondary malignancy originating within the SIP. It is considered primary if there is no additional mucosal condition or secondary if it occurs in conjunction with an additional condition, which is estimated to occur in about 9% of cases. [3] SIP is usually not a primary lacrimal drainage system tumor but a primary tumor of the tissue surrounding the lacrimal drainage system and spreads to the lacrimal sac via secondary extension. It is exceedingly rare for SIP to originate from the lacrimal sac, having been reported only 7 times in the past 30 years. [7]

To date, the treatment for SIP includes surgery and surgery combined with radiotherapy for SIP-associated squamous cell carcinoma (SIP/SCC). Currently, the majority of surgeons prefer endoscopic methods to traditional external approaches, due to similar success rates, less trauma and no facial scars. However, the common view is that SIP recurrence is due to inadequate removal during the first surgery [8]. Therefore, preoperative evaluation as well as postoperative follow up is crucial. The strength of this case report is the thorough diagnostic work-up in such a paucisymptomatic patient. The main limitation is that it was not possible to assess the exact moment and duration of onset of symptoms.

Conclusions

The present case points out the importance of including mass in differential diagnosis of nasolacrimal duct obstruction when epiphora is the only manifest sign.

Masses are uncommon but not a rare cause of nasolacrimal duct obstruction. Surgical teams performing high numbers of dacryocystorhinostomy procedures should be aware of such pathology and follow a systematic multidisciplinary approach. This is essential since neoplastic secondary acquired nasolacrimal duct obstructions (neoplastic SANDOs) can be easily mistaken for benign processes such as primary acquired nasolacrimal duct obstructions (PANDOs). [9]

We subject all candidates for DCR surgery to a preoperative dacryo-CT scan to exclude facial bone anomalies and concomitant tear duct stenosis. We believe that dacryo-CT scan is the most useful exam for evaluation of the lacrimal drainage system, together with nasal cavities and sinuses. This case demonstrates that bone destruction with intraorbital and intracranial extension are possible severe complications. Whenever a malignant tumor expands beyond the lacrimal sac, it should be treated according to the degree of histological pattern and extension within the orbital soft tissue. The major cause for recurrence of the disease, or rather failure to cure it, is in case of incomplete resection of the tumor or insufficient cleanliness of the surgical margin. [9]

Every attempt should be made to remove the whole tumor during primary surgery. To obtain neoplastic radicality, we proceeded with orbital exenteration combined with lateral rhinotomy. It is intuitive that radical surgery can be most easily performed in the initial stages of the disease.

Our patient reported having just persistent epiphora for at least 1 years prior to presentation of nasal symptoms, when she also noted that a right medial canthal swelling had recently arisen. Therefore, unilateral persistent epiphora in patient at risk must be taken seriously. This is the primary take-away lesson. Patient perspective on treatment was concerning the unexpected severity of the disease if compared to the relatively mild symptoms.

Recurrence rate for inverted papilloma has been described in up to 78% in the literature and some risk factors for recurrence were identified, such as tobacco exposure, size of tumor, high hyperkeratosis, squamous hyperplasia, increased number of mitosis, HPV positivity and tumor location. In most cases, recurrence occurs in the first three years after surgery, although, there are reports of recurrence after 10 years. [10] So, every attempt should be made to remove the whole tumor during primary surgery and long-term follow-up is required in order to rule out recurrences. In our case both Ki-67 and HPV were negative and our five-year follow-up did not reveal any recurrence. Because of the rarity of the tumors, we believe it is advisable that such cases be managed in a multidisciplinary setting through a systematic approach.

Declarations

I confirmed we gained ethics approval and consent to participate. I confirmed we gained consent for publication. I confirmed we have no competing interests. I confirmed we received no funding. The authors have no financial disclosure.

“Written informed consent was obtained from the patient for publication of this case report and any accompanying images. A copy of the written consent is available for review by the Editor-in-Chief of this journal.”

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