Case Report

Severe epistaxis in pregnancy due to nasal pyogenic granuloma: A case report

Johannas Mohd Yusof, MBBCh a,*, Azwarizan Abd Halim, MD ORL-HNS b and Aneeza Khairiyah Wan Hamizan, PhD a

a Department of Otorhinolaryngology-Head and Neck Surgery, Hospital Canselor Tuanku Muhriz UKM, Kuala Lumpur, Malaysia
b Otorhinolaryngology Department, Hospital Tuanku Jaafar, Seremban, Malaysia

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Abstract

Mild to severe epistaxis is common in pregnancy and often results from increased vascularity of the nasal mucosa and hormonal changes. Symptoms may occur in the absence of an obvious local cause or any systemic disorder; however, thorough otolaryngological (i.e., “ENT”) evaluation is always warranted. Pyogenic granuloma or lobular capillary haemangioma is a benign fibrovascular proliferative tumour that is commonly found on the face, fingers, lips, and nasal mucosa. Pregnancy-induced pyogenic granuloma is not an uncommon entity and may result in torrential epistaxis if untreated. Managing a case of severe epistaxis during pregnancy usually requires multidisciplinary management. The authors present a case of severe epistaxis in pregnancy that necessitated examination of the nasal cavity under general anaesthesia. Intraoperative findings showed a bluish-red mass occupying the patient’s right maxillary sinus. Bleeding was arrested and complete haemostasis was achieved. The nasal pyogenic granuloma completely resolved in the post-partum period.

Keywords: Capillary haemangioma; Epistaxis; Multidisciplinary management; Nasal pyogenic granuloma; Pregnancy

Introduction

Lobular capillary haemangioma or pyogenic granuloma is a benign fibrovascular proliferative tumour occurring in both...
Children and adults. Occurrence during pregnancy is reported to be approximately 5%. Based on a 10-year retrospective analysis of 155 cases by Koo et al., the predominant site for pyogenic granuloma is the face (30.3%), followed by the fingers (22.5%) and lip and oral cavity (10%). Nasal mucosa involvement is fairly common and has been reported to be as low as 1% and as high as 33% of cases. Lopez et al. reported that the most common site for sinonasal pyogenic granuloma is the nasal septum, followed by the inferior turbinates. The occurrence of maxillary sinus pyogenic granuloma, however, has scarcely been reported.

Pregnancy is associated with up to 15% of nasal pyogenic granulomas, while a history of trauma was reported in 12% of these cases. Etiological factors for nasal pyogenic granuloma include increased or altered hormone activity (40%), previous injury to the nasal cavity (18%), and potential idiopathic disease in approximately 23% of cases. Some studies have also reported an increased frequency of pyogenic granuloma in patients with pre-existing skin conditions, including dermatitis, or vascular malformations such as hemangioma.

Case report

A 26-year-old primigravida at 35 weeks’ gestation presented with persistent unilateral epistaxis, soaking more than one-half of her handkerchief. She claims to have experienced intermittent unprovoked epistaxis, which usually resolved spontaneously since entering her third trimester. She experienced rhinitis symptoms but denied any history of local trauma and no other bleeding tendencies. She reported a surgical history of right nasal polypectomy approximately 5 years previously.

Physical examination showed slow active oozing from the patient’s right nostril; however, she was hemodynamically stable. First aid Trotter’s manoeuvre with ice compression was attempted at the emergency department, and she was transferred to the authors’ clinic for evaluation. Her epistaxis resolved after ice compression, thus enabling rigid nasal endoscopy to be performed, which showed clots over the lateral nasal wall. Initial complete blood count showed a haemoglobin level of 12.1 g/dl and platelet count of 206 x 10^9/l, with normal coagulation profile. She was admitted for observation; however, after 2 h in the ward, she experienced another active episode that was not resolved with compression. Accordingly, anterior nasal packing using Merocel (Medtronic Inc., Minneapolis, MN, USA) was performed and empirical intravenous (IV) amoxicillin clavulanate (Augmentin, GlaxoSmithKline, Brentford, United Kingdom) was started.

The packing was removed after 48 h; however, the patient experienced another episode of severe epistaxis that required both anterior and posterior nasal packing using a Foley’s catheter. At this point, the patient experienced stage 1 hypovolemic shock and was resuscitated using 0.9% normal saline (IV, 10 ml/kg) over a 30 min period. After discussion between the obstetrician and patient, a decision was made for examination under anaesthesia to identify the source of epistaxis, and to secure haemostasis with the possibility of endoscopic sphenopalatine artery ligation. Differentials at this point included nasal pyogenic granuloma, bleeding nasal polyp, sinonasal tumour, and vascular malformation.

Intraoperatively, the nasal packing was removed, and rigid nasal endoscopy was performed. A fleshy reddish mass, measuring approximately 1 x 1 cm, was seen arising from the right lateral nasal wall adjacent to a widened maxillary antrum (Figure 1). On removal, the mass measured 1.0 x 1.5 cm and was sent for histopathological examination; however, it was reported to only be blood clots. A widened middle meatal antrostomy was evident, and a smooth blush-red mass with active oozing arising from the anterior wall of the maxillary antrum was noted (Figure 2). The bleeding points were cauterized using suction diathermy, and an absorbable haemostat (Surgicel, Ethicon/Johnson & Johnson, Somerville, New Jersey, USA) was placed over the tissue to achieve haemostasis. Blood loss during the operation was estimated to be approximately 250 ml.
The patient recovered well during the immediate post-operative period. She experienced no recurrent epistaxis and was subsequently discharged at day 2 postoperatively. She was followed up 2 weeks later and subsequently at 1 month post parturition. During her last follow up, she denied any episodes of epistaxis or nasal obstruction. Rigid nasal endoscopy showed complete resolution of the previously observed lesion within the right maxillary sinus.

Discussion

Pregnancy granuloma or granuloma gravidarum are among other terms for lobular capillary haemangioma occurring in pregnancy. It is the most common vascular lesion of the nasal cavity. In one of the largest case series investigating lobular capillary haemangioma, the lesion occurs more in women, with a ratio of 2:1, and at a mean age of 45 years. The majority of patients present with epistaxis followed by nasal obstruction.

The aetiology of lobular capillary haemangioma is unclear; however, theories associated with hormonal components have been proposed because the lesion is commonly encountered in pregnant patients and those taking oral contraceptives. History of nasal injury is also a possible aetiology, ranging from 11 to 15% of nasal pyogenic granulomas. Some case reports have described the occurrence of nasal pyogenic granuloma after nasal surgery and nasal packing, which are believed to trigger an inflammatory reaction in response to the exposed mucosal surfaces. It is usual for it to occur years after trauma, given that most reported cases occur within 1–2 months of injury; however, it is possible that our patient had been asymptomatic up until her first pregnancy due to hormonal changes, which have also been attributed to the pathophysiology of sinonasal pyogenic granuloma.

Although the molecular mechanism of lobular capillary haemangioma regression post parturition is unclear, it has been proposed that it is due to a decrease in vascular endothelial growth factor levels post parturition. Hence, it has been suggested that excision is indicated only in cases that fail to involute. Surgical excision is the preferred treatment for lobular capillary haemangioma of the nasal cavity. Other methods include electrocautery, laser therapy, and embolization. Recurrence can be as high as 40%, especially for lesions managed using cautery. Managing epistaxis in pregnancy can be difficult because some treatments may be contraindicated. Maternal safety needs to be prioritized with conservative measures being the first-line treatment(s). As described by Crunkhorn et al., chemical cautery with silver nitrate is safe in mild cases, as is anterior and posterior nasal packing; however, the use of bismuth iodiform paraffin paste-soaked ribbon gauze for nasal packing is contraindicated. If epistaxis is severe and conservative management fails, surgical intervention, such as vessel ligation, is feasible with consideration of the risks of general anaesthesia in pregnancy. Cocaine-based nasal preparations should be avoided due to well-known risks to the fetus.

In the present case, the patient underwent examination under anaesthesia due to severe epistaxis and no preoperative radiological investigation(s) could be performed. A clinical decision was made to manage the lesion conservatively to achieve haemostasis with the possibility of second-look surgery post-partum. This was because there were no computed tomography imaging data to guide in terms of anatomical relations of the lesion to other important structures. There was also suspicion of it being a vascular lesion; hence, we did not want to risk profuse bleeding with subsequent harm to both patient and foetus. The diagnosis was made only in retrospect, after involution of the lesion post-partum.

In conclusion, epistaxis in pregnancy can be severe due to hormonal changes, with subsequent increase in nasal vascularity. Persistent severe epistaxis in pregnancy that is refractory to conservative treatment necessitates surgical treatment to achieve haemostasis. Although nasal pyogenic granuloma is a common benign cause of epistaxis in pregnancy and usually regresses post-partum, all pregnant patients with persistent epistaxis should be referred for otolaryngological (i.e., “ENT”) assessment to prevent significant morbidity.

Recommendation

The authors stress the importance of complete ENT assessment and evaluation in a pregnant patient presenting with recurrent or persistent epistaxis to the primary health care or obstetric physician.

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Conflict of interest

The authors have no conflict of interest to declare.

Ethical approval

No personal details or identifying information are included in this article. Consent was provided by the patient for the case report to be written and published. Ethics approval was not obtained from the authors’ Clinical Research Center because it was not required for a case report.

Authors contributions

JMY collected necessary case information and wrote the original and final draft. AAH conceptualized the case report and its aims and reviewed the initial draft. AKWH provided critical review and editing of the case report. All authors have critically reviewed and approved the final draft and are responsible for the content and similarity index of the manuscript.

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