Congenital bilateral adult choanal atresia undiagnosed until the second decade: How we did it

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ABSTRACT

Background: Bilateral congenital choanal atresia that presents in adulthood is rare. There are only eight reported cases in the literature.

Method: We present a ninth case of adult bilateral choanal atresia diagnosed at the age of 20 years. Can a bilateral choanal atresia present so late in life when such a condition is considered incompatible with life?

Results and conclusion: Clinical details, diagnosis, and surgical steps with a clear depiction of photographs and comparison with all other previously reported cases, which can help novel otolaryngologists in their clinical practice are discussed. (Allergy Rhinol 7:e82–e84, 2016; doi: 10.2500/ar.2016.7.0155)

Congenital choanal atresia is a condition in which the nasal cavity fails to communicate with the nasopharynx. This condition almost always presents in neonates, with history of episodes of cyclical cyanosis and respiratory distress that worsens during feeding and is relieved by crying. Up to two-thirds of cases of choanal atresia are unilateral, and bilateral cases are commonly associated with other congenital anomalies. It is not uncommon for unilateral disease to present later in life, whereas bilateral disease is considered to be incompatible with life and is rarely seen in adults. An extensive search for bilateral congenital choanal atresia in adults, revealed, to our knowledge, only eight reported cases to date.1–8 We present a ninth case of congenital bilateral choanal atresia in a 20-year-old woman and describe the clinical features, evaluation, and surgical steps to help novel otolaryngologists in their practice.

CASE REPORT

A 20-year-old woman with a poor socioeconomic status presented to the outpatient clinic with chief concerns of nasal obstruction, nasal discharge, snoring, anosmia, and mouth breathing since childhood. Her birth history was uneventful; there was a history of episodes of cyanosis in early childhood, with recovery on crying. There was a history of frequent hospital admissions for recurrent pneumonia in childhood. There was no history of trauma to the nose. Results of a clinical examination revealed long face with overcrowded incisors, a high-arched palate, and hypoplasia of the maxilla. There were no other congenital anomalies.

On nasal endoscopy, both nasal cavities were narrow, with absent posterior choanae. Therefore, a diagnosis of bilateral choanal atresia was made (Fig. 1). Computerized tomography of the nose and paranasal sinus showed a thick hypodense sheet of soft tissue completely obliterating the bilateral choanae. There was bowing of the bilateral lateral nasal walls, with thickening of the vomer (Fig. 2).

An endoscopic choanoplasty was performed in which the membranous atresia was removed by using a microdebrider, and drilling of the vomer and medial pterygoid plates was performed (Fig. 3). A bilateral nasal pack was kept for 5 days. No stenting was done. Thorough nasal douching was advised for the next 2 weeks. At the 3-month follow-up, there were well-patent posterior choanae (Fig. 4). Informed written consent was given by the patient for publication of her case and accompanying images.

DISCUSSION

Choanal atresia has an incidence of 1 in 5000 to 1 in 8000 live births.9 Fifty percent of all patients with choanal atresia and up to 75% of patients with bilateral disease have other associated congenital anomalies.10 Most common syndromes associated include CHARGE syndrome, Crouzon syndrome, and craniosynostosis. The incidence of mixed atresia is more common than pure membranous and bony atresias11 as also seen in our case.

Unilateral disease presents most commonly in the later part of life, with nasal discharge and obstruction. Because neonates are obligate nasal breathers, bilateral disease most often presents in the perinatal period, with attacks of cyanosis and apneic attacks.12 However, in the present
case, the parents did not find cyanotic spells, but there were multiple hospital admissions in the patient’s childhood. Our patient presented with symptoms of nasal obstruction, discharge, snoring, and mouth breathing, as also noted by Tinoco et al.¹

Nasal endoscopy in our case revealed narrow nasal cavities, thickening of the posterior septum (vomer), medially shrunken lateral nasal walls at the posterior choanae (pterygoid plates), as also noted by other investigators.¹³¹⁴ Noncontrast computerized tomography of nose and paranasal sinuses helps in assessing the thickness of the atresia and vomer and the position of pterygoid plates, and to distinguish between membranous and bony atresia; it also helps to know the status of the paranasal sinuses.

**Figure 1.** Nasal endoscopic view of posterior choanae, depicting complete atresia.

**Figure 2.** Noncontrast computerized tomography, axial section at the level of posterior choanae, depicting mixed atresia.

**Figure 3.** Immediate postoperative view of right posterior choana after transnasal endoscopic choanoplasty.

**Figure 4.** A 1-year follow-up endoscopic photograph, depicting well-epithelized patent posterior choanae.
Transnasal endoscopic choanoplasty was attempted in our patient. The membranous atresia was initially cleared by using a microdebrider. Bony atresia was drilled by using a 2.5-mm cutting burr under direct visualization.\(^2\) The thickened vomerine bone and medial pterygoid plates were drilled until proper visualization of the Eustachian tube opening. The role of a stent is controversial. Some investigators advocate its supremacy.\(^2\) Aksoy et al.\(^3\) used mitomycin C after choanoplasty. In our case, neither of these adjuvants were used. In the postoperative period, the nasal pack was removed after 5 days, and the patient started with nasal douching for 2 weeks. Our patient was followed up for 3 months, and nasal endoscopy revealed well-patent posterior. Of the eight documented cases of bilateral choanal atresia,\(^1,3,8,14\) in the literature, only one patient had restenosis, which required revision surgery.\(^4\)

CONCLUSION

Bilateral choanal atresia is considered incompatible with life. Early surgical intervention in neonatal period is necessary for survival. Adult bilateral choanal atresia is a rare entity. Bilateral nasal obstruction, recurrent sinusitis, snoring, and mouth breathing are usual symptoms at the time of presentation. Nasal endoscopy and preoperative computerized tomography helps in planning surgery. Endoscopic transnasal choanoplasty is the criterion standard treatment. The exact role of postoperative stenting and the use of mitomycin C are controversial.

REFERENCES

1. Tinoco P, Pereira JCO, Caldas Lourenço FR et al. Bilateral choanal atresia in a 34 year old patients. Int Arch Otorhinolaryngol 14:481–484, 2010.
2. Panda NK, Sunhadri S, and Ghosh S. Bilateral choanal atresia in an adult: Is it compatible with life? J Otorhinolaryngol 118:244–245, 2004.
3. Aksoy F, Demirhan H, Yildirim YS, and Ozturan O. Bilateral choanal atresia in an adult: Management with mitomycin C and without stents: A case report. Cases J 2:9307, 2009. Available online at http://www.casesjournal.com/content/2/1/9307.
4. El-Sawy H, Siddiq MA, and Anbarasu A. Bilateral choanal atresia and paranasal sinus hypoplasia in adult patient with hypogammaglobulinemia. Eur Arch Otorhinolaryngol 263:1136–1138, 2006.
5. Iseh KR, Nasir A, and Noma U. Adult bilateral bony choanal atresia with right ethmoidal mucocele, right chronic dacryocystitis and dacryocystocele. Nat J Otorhinolaryngol Head Neck Surg 2:23–24, 2014.
6. Yasar H, and Ozkul MH. Bilateral congenital choanal atresia in a 51-year-old woman. Am J Rhinol 21:716–718, 2007.
7. Tatar EC, Ozdek A, Akcan F, and Korkmaz H. Bilateral congenital choanal atresia encountered in late adulthood. J Laryngol Otol 126:949–951, 2012.
8. Chaudhary N, Jain A, Kapoor R, and Motwani G. Bilateral complete choanal atresia in an adult woman: Managed with nasal endoscopes. J Indian Med Assoc 108:109–110, 2010.
9. Pirsig W. Surgery of choanal atresia in infants and children: Historical notes and updated review. Int J Pediatr Otorhinolaryngol 11:153–170, 1986.
10. Brown K, Rodriguez K, and Brown OE. Congenital Malformations of the Nose. 4th ed. Philadelphia: Elsevier Mosby, 4099–4109, 2005.
11. Brown OE, Pownell P, and Manning SC. Choanal atresia: A new anatomic classification and clinical management applications. Laryngoscope 106:97–101, 1996.
12. Elluru RG and Wootten CT. Congenital malformations of nose. Cummings Text Book of Otolaryngology and Head Neck Surgery, 5th ed., Vol. 3, 2010.
13. Harner SG, McDonald TJ, and Reese DF. The anatomy of congenital choanal atresia. Otolaryngol Head Neck Surg 89:7–9, 1981.
14. Brown OE, Smith T, Armstrong E, and Grundfast K. The evaluation of choanal atresia by computed tomography. Int J Pediatr Otorhinolaryngol 12:85–98, 1986.