Melanotic Oncocytic Metaplasia of the Nasopharynx in the Patient with Suspicious Hemoptysis: Case Report

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Melanotic oncocytic metaplasia (MOM) in the nasopharyngeal space is a very rare entity. Only 35 cases have been reported in the English literature, and most patients were East Asian males between 60 and 70 years of age. MOM presents as a brown or black lesion with slight elevation of the mucosa. These lesions are benign and defined as cellular enlargement with eosinophilic granular melanin-pigmented cytoplasm caused by mitochondrial accumulation. However, such presentation can lead physicians to misjudge MOM as a malignant lesion. Recently, we experienced a case of MOM of the nasopharynx. A 58-year-old woman was admitted to the internal medicine department with small-volume hemoptysis and referred to the ENT department for evaluation. She was a regular smoker without any medical history. Sinus endoscopy showed black pigmented lesions on both the torus tubaris and left posterior tonsillar pillar, with low bleeding risk. Excisional biopsy of the lesion was performed, and oncocytic metaplasia was confirmed pathologically. Hemoptysis showed spontaneous remission and no recurrence or other symptoms over 12 months of follow up. Melanotic oncocytic metaplasia in the nasopharynx should be clinically recognized to avoid misdiagnosis as a malignancy like melanoma.

KEY WORDS: Adenoma, Oxyphillic · Nasopharynx · Hemoptysis.

INTRODUCTION

First melanotic oncocytic metaplasia (MOM) of the nasopharynx was reported in 1995.¹ MOM can be misdiagnosed as early nasopharyngeal carcinoma or melanoma due to its macroscopic appearance. However, MOM is clearly distinct from such diseases histologically. MOM comprises both oncocytic metaplasia and melanin pigmentation of the epithelium in the same gland. Oncocytes are big epithelial cells with an aplenty, deeply eosinophilic granular cytoplasm and a small, round, dark colored nucleus. Melanocytes found to coexist in MOM, is considered to be the source of melanin.² No atypia or mitotic activity found in MOM. Fontana-Masson staining is positive and highlights several dendritic melanocytes between the metaplastic and the surface epithelial cells.³ MOM is thought to be an age-related change since its incidence increases with age, and predominantly occurs in Asian population.⁴

MOM of the nasopharynx is a very rare lesion, and only 35 cases were reported in English literature. Most lesions were asymptomatic and incidentally discovered during physical examination.

We present here a case of MOM of the nasopharynx, found in a patient with suspicious hemoptysis.

CASE REPORT

A 58-year-old woman was admitted to internal medicine department, presenting small volume hemoptysis for 5 days. She was a regular smoker (10/day × 20 years) without any
medical history. Patient admitted and undergone computed tomography scan of lung. Since the radiologic evaluation at the internal medicine department showed no clinical significance except for minor bronchopneumonia, consultation to ENT department was proceed for further evaluation of other possible causes. Sinus endoscopy revealed black pig-

![Fig. 1. Endoscopic findings. A: Endoscopic finding of melanotic oncocytic metaplasia on left anterior pharyngeal pillar (white arrow). B: Endoscopic finding of melanotic oncocytic metaplasia on both torus tubarius (white dotted arrows).](image)

![Fig. 2. Pathologic results of the case. A: Microscopic finding (×100) shows deeply homogeneous eosinophilic epithelial cells, noted with lumen formation. B: Scattered melanin pigment granules are found in eosinophilic cytoplasm (×400).](image)

![Fig. 3. Endoscopic findings after 12 months. A: Rt torus tubarius. B: Lt torus tubarius.](image)
mented lesions on both torus tubaris and left posterior tonsillar pillar (Fig. 1), but bleeding from the lesion was unseen. Sinus mucosa and anatomy was intact otherwise. Though hemoptysis stopped spontaneously during the admission period, surgical biopsy of the lesion was performed to rule out possible malignancy. Five specimens were sent for he-

| Case | Sex/age | Site | Chief complaint for visiting clinic | Number of lesions | Smoking history | Author/year |
|------|---------|------|------------------------------------|------------------|----------------|-------------|
| 1    | M/67    | EO   | Otitis media                       | Single           | Unknown        | Shek TW, 1995 |
| 2    | M/63    | EO   | Tinnitus                           | Single           | Unknown        | Shek TW, 1995 |
| 3    | M/70    | Bilateral EO | Tinnitus                      | Multiple         | Unknown        | Xue WC, 1999 |
| 4    | M/64    | Bilateral EO, left suprapharynx, and nasal cavity | Throat discomfort | Multiple         | Unknown        | Hirakawa E, 1999 |
| 5    | M/62    | Left EO and bilateral torus tubaris | Discomfort in Lt ear | Multiple         | Unknown        | Takano K, 2004 |
| 6    | M/69    | Left nasopharynx | Foreign body swallowing         | Multiple         | Unknown        | Kunihara K, 1997 |
| 7    | M/80    | Right nasal cavity and pharynx | Hoarseness             | Multiple         | 10/day × 50 yr | Sakaki M, 2004 |
| 8    | M/69    | Left EO | Hoarseness                         | Single           | 40/day × 60 yr | Sakaki M, 2004 |
| 9    | M/74    | Left nasopharynx | Rhinorrhea                 | Single           | Unknown        | Sakaki M, 2004 |
| 10   | F/74    | Right EO | Throat discomfort                | Multiple         | Unknown        | Sakaki M, 2004 |
| 11   | M/68    | Nasopharynx | None                             | Single           | Unknown        | Sakaki M, 2004 |
| 12   | M/65    | Right EO | Hemoptysis                        | Single           | Unknown        | Sakaki M, 2004 |
| 13   | M/63    | Left EO | Epistaxis                          | Single           | Unknown        | Sakaki M, 2004 |
| 14   | M/-     | Bilateral nasopharynx | None(examined to find origin site of metastatic carcinoma) | Multiple         | Unknown        | Liao C, 2005 |
| 15   | M/79    | Right EO | Otitis media                       | Single           | 20/day × 40 yr | Lui PC, 2004 |
| 16   | M/58    | Bilateral nasopharynx | Epistaxis                  | Multiple         | Unknown        | Li Y, 2010 |
| 17   | M/73    | Bilateral torus tubaris | Nasal congestion            | Multiple         | Non-smoker     | Kondo, 2010 |
| 18   | M/72    | Bilateral torus tubaris | Headache and hearing impairment | Multiple         | 40/day × 50 yr | JY Na, 2012 |
| 19   | M/71    | Left torus tubaris and soft palate | Hoarseness              | Multiple         | 20/day × 40 yr | JY Na, 2012 |
| 20   | M/51    | Right torus tubaris | Tongue pain                 | Multiple         | Unknown        | JY Na, 2012 |
| 21   | M/63    | Nasopharynx | Epistaxis                  | Multiple         | 40/day × 40 yr | Chang, 2014 |
| 22   | M/57    | Right torus tubaris | None(examined to find origin site of metastatic carcinoma) | Multiple         | 30/day × 40 yr | Shogo Tajima, 2015 |
| 23   | F/70    | EO | Routine gastrointestinal endoscopy | Multiple         | Unknown        | Uehara K, 2015 |
| 24   | F/61    | Left EO | Hoarseness                     | Single           | Unknown        | Uehara K, 2015 |
| 25   | M/74    | Bilateral nasopharynx | Hoarseness                 | Multiple         | Unknown        | Uehara K, 2015 |
| 26   | M/57    | Right nasopharynx | Nasal obstruction            | Single           | Smoker         | Li J, 2019 |
| 27   | M/61    | Left nasopharynx | Neck mass                    | Single           | Smoker         | Li J, 2019 |
| 28   | M/69    | Left nasopharynx | Rhinorrhea                   | Multiple         | Smoker         | Li J, 2019 |
| 29   | M/56    | Left EO | Epistaxis                      | Single           | Smoker         | Li J, 2019 |
| 30   | M/58    | Right nasopharynx | Facial palsy               | Single           | Smoker         | Li J, 2019 |
| 31   | M/52    | Left EO | Tinnitus                       | Multiple         | Smoker         | Li J, 2019 |
| 32   | F/77    | Left nasopharynx | Hoarseness                  | Single           | Smoker         | Li J, 2019 |
| 33   | M/59    | Left nasopharynx | Nasal obstruction         | Single           | Smoker         | Li J, 2019 |
| 34   | M/59    | Left EO | Tinnitus                      | Single           | Smoker         | Li J, 2019 |
| 35   | M/75    | Right EO | Epistaxis                  | Multiple         | 30/day × 50 yr | Chen HY, 2020 |
| 36   | F/53    | Left posterior tonsillar pillar, both torus tubaris | Hemoptysis           | Multiple         | 10/day × 20 yr | Present case, 2020 |

EO: Eustachian opening, M: male, F: Female
matoxylin and eosin (H&E) staining. Microscopically, homogenous oncocytic cells with melanin pigment granules in their cytoplasm were found, and melanotic oncocytic metaplasia was pathologically confirmed (Fig. 2). Hemoptysis stopped spontaneously. The patient discharged without complications and didn’t presented hemoptysis or other symptoms during 12 months follow up. Endoscopic finding showed no further growth or advancement of previous lesion (Fig. 3).

**DISCUSSION**

Oncocytic change found in the upper respiratory tract is an unusual finding. Melanotic variation of oncocytic metaplasia is an extremely anomalous condition, presented as a small, brown to black mucosal lesion.\(^1\) To our knowledge, only 35 cases have been reported in the English literature, and our case is the 36th (Table 1). All reported cases were elderly Asian, supporting the possibility of ethnic back-ground as a predisposing factor of MOM. MOM predominantly occurred in males (31/36), and most of patients were long time smoker (18/19), supporting the hypothesis that smoking might be a predisposing factor for MOM. The male predominance of patients supports this hypothesis also, since male smokers are more frequent among Asian population. 16 cases showed single lesion while other 20 cases presented multiple lesions in nasopharynx. The presented patient was an Asian female with multiple lesions, and smoking history was relatively short compared to other 17 previously reported cases with smoking history.

Pathological findings of the patient in H&E stain showed oncocytic metaplasia and scattered melanin pigment granules. Atypia or mitotic figures were not found. Oncocytic cells are epithelial cells, characterized by abundant eosinophilic cytoplasm. Yet, oncocytic metaplasia is uncommon finding in upper respiratory tract. The exact origin of the melanin pigment in MOM is still in question. However, previous studies demonstrated that Fontana-Masson stain, S-100 protein immunostain and human melanoma black-45 immunostain revealed dendritic melanocytes stretching through the epithelial cells of the glands in MOM,\(^6\) supporting that melanin pigment may be derived from the adjoining melanocytes through their stretching dendrites.\(^4\)

Most of the cases were found incidentally with other symptoms such as tinnitus, hoarseness, epistaxis, rhinorrhea, and hemoptysis. Some symptoms, such as ear discomfort at the same side of MOM near eustachian opening, seemed to be related with the lesion, while other symptoms like hoarseness seemed almost independent from the lesion.

It is unclear whether MOM can cause hemoptysis or not. One case with hemoptysis was reported previously,\(^9\) but it’s clinical course or relation between the lesion was not described. Three cases with epistaxis had been reported also. During the endoscopic biopsy of presented case, the lesions didn’t show hemorrhagic features, supporting irrelevance between hemoptysis and the lesion. Minor bronchopneumonia could have been the origin of hemoptysis in this case, but was not clearly identified.

Clinically, carcinoma or melanoma may show similar appearance with MOM during the endoscopic examination. Differential diagnosis between those entities are straightforward histologically, so physicians encountered such lesions should try biopsy. Endoscopic findings isn’t sufficient to distinguish between MOM and nasopharyngeal malignancy since nasopharyngeal malignancy may shows various forms including submucosal lesions.

It is known that MOM follows a benign clinical path. Simple excision is a suitable treatment for MOM.\(^2\) No abnormal bleeding or surgical difficulty was found in operation of this case. Since no recurrence or advancement of the lesion has been reported in the literatures, diagnosed MOM can be observed without any further treatment.

In conclusion, we report the 36th cases of melanotic oncocytic metaplasia in the nasopharynx. As a benign imitator of malignant lesion, MOM should be always taken into consideration in examination of nasopharynx.

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