Primary intratracheal neurilemmoma in a 10-year-old girl: A case report

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Abstract

BACKGROUND
Tracheal tumors are relatively rare in adults and uncommon in children. Tracheal neurilemmoma is a rare condition in adults that usually affects middle-aged people, but it can also occur in children. Because the clinical presentation is nonspecific and insidious, diagnosis is often delayed. The most common symptoms in these patients are stridor or wheezing (especially positional) and cough. A few patients are misdiagnosed and mistakenly treated for asthma.

CASE SUMMARY
A 10-year-old girl was admitted to our unit with a 2-mo history of recurrent cough, dyspnea, and tachypnea. Her condition was more severe after exercise. Her symptoms progressed despite treatment with inhaled fluticasone/salmeterol. Flexible electronic laryngoscopy showed a red, smooth, and round mushroom-shaped mass in the trachea, about 1 cm below the vocal cords. The surface of the
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mass was covered with several small and discontinuous blood vessels. About 90% of the tracheal lumen was occupied by the mass. A multidisciplinary operation was performed. The surgically resected mass was diagnosed as benign neurilemmoma by immunohistochemical analysis.

**CONCLUSION**

Intratracheal neurilemmoma is fairly rare in children. The main symptoms include coughing, wheezing, and dyspnea. The tumor’s size, location, and degree of intratracheal and extratracheal invasion can be measured by chest computed tomography. The main treatment strategies used for tracheal neurilemmoma are surgical resection and endoscopic excision. Long-term follow-up is warranted for the evaluation of outcomes and complications.

**Key Words:** Airway obstruction; Trachea; Neurilemmoma; Therapeutic bronchoscopy; Child; Case report

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**Core Tip:** Tracheal neurilemmoma is a rare condition in children. Because the clinical presentation is nonspecific and insidious, diagnosis is often delayed. This case report describes a 10-year-old girl with tracheal neurilemmoma.

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**INTRODUCTION**

Tracheal tumors are relatively rare in adults and uncommon in children[1]. About two-thirds of such tumors are malignant or moderately malignant, and the remaining one-third are benign lesions[2]. Tracheal neurilemmoma, also known as schwannoma, is a rare condition in adults that usually affects middle-aged people; however, it can also occur in children[3]. Common symptoms of this condition are stridor or wheezing and coughing. Some patients are misdiagnosed and mistakenly treated for asthma. This report describes a case involving a 10-year-old girl with tracheal neurilemmoma.

**CASE PRESENTATION**

**Chief complaints**

A 2-mo history of recurrent cough, dyspnea, and tachypnea.

**History of present illness**

A 10-year-old girl was admitted to our unit with a 2-mo history of recurrent cough, dyspnea, and tachypnea. Her condition was more severe after exercise. She had been hospitalized twice in a local unit and diagnosed with bronchitis and asthma. Her symptoms progressed despite treatment with inhaled fluticasone/salmeterol.

**History of past illness**

The patient had been previously healthy, and she denied a history of asthma and seasonal allergic rhinitis.

**Personal and family history**

The patient had no relevant family history.
Physical examination
On admission, the patient was dyspneic and characterized by biphasic stridor with a percutaneous saturation of 95% on room air. An examination showed a body temperature of 36.7 ºC, a pulse rate of 92 beats/min, a respiration rate of 34 breaths/min, and a blood pressure of 107/74 mmHg.

Laboratory examinations
The patient’s leukocyte count was 6.07 × 10^9/L (reference range: 4.0-12.0 × 10^9/L). Her lymphocyte count was 3.09 × 10^9/L (reference range: 0.7-4.9 × 10^9/L). An immune function test showed that her immunoglobulins were within the reference range.

Imaging examinations
Contrast-enhanced computed tomography (CT) revealed an obviously enhanced mass, occluding about 90% of the tracheal lumen (Figure 1A and B). Under conscious and local anesthesia, flexible electronic laryngoscopy showed a red, smooth, and round mushroom-shaped mass. The surface of the mass was covered with several small and discontinuous blood vessels. The base of the mass was in the trachea, about 1 cm below the vocal cords at the 4- to 7-o’clock position (Figure 2A).

Pulmonary function examinations indicated that the patient, who had exhibited no marked response to bronchodilators, had severe inspiratory obstructive ventilation dysfunction.

FINAL DIAGNOSIS
According to the pathological and immunohistochemical examination findings, the patient was diagnosed with tracheal neurilemmoma.

TREATMENT
The multidisciplinary consensus met the performance of endoscopy for a definitive diagnosis and determined prognosis and guide management. After the patient’s parents had been fully notified of surgical options and possible complications associated with the various surgical modalities, they agreed to endoscopic tumor resection. There was a rigid bronchoscope available as well as a low-temperature plasma therapy device, argon plasma coagulation device, cryotherapy device, and holmium laser if necessary. The patient’s parents also consented that, in case of failed endoscopic tumor resection, surgical resection would be conducted instantly.

Otolaryngologists, pulmonologists, anesthesiologists, thoracic surgeons, and doctors of the pediatric intensive care department were on-call in the operating room, ensuring the patient’s safety. The loading dose of dexamethasone was 1 µg/kg, and the maintenance rate was 0.3 µg/kg/h. The patient was maintained with 3% sevoflurane inhalation, remifentanil at 0.2 µg/kg/min, intravenous methylprednisolone at 30 mg, intravenous atropine at 0.25 mg, and intravenous lidocaine at 25 mg. After deep sedation and spontaneous breathing were achieved, 10 mg of etomidate was added to deepen the anesthesia. A flexible bronchoscope (FB) with a 2.8 mm outer diameter (Olympus XP260; Olympus Corp., Tokyo, Japan) was inserted into the trachea over the mass and used to guide intubation of a 4.5 mm endotracheal tube. The 4.5-mm tube could not prevent blood from flowing into the lower airway; therefore, we decided to perform a tracheotomy below the mass. The anesthesia ventilator controlled the patient’s breathing through a tracheotomy cannula. After extubation of the endotracheal tube, a laryngoscope was placed in the vallecula, exposing the supraglottic and glottic larynx. An FB with a 4.9 mm outer diameter (Olympus 260; Olympus Corp.) was then passed into the subglottic region. A flexible electrocautery snare with a 1.9 mm outer diameter (Olympus CD-6C-1; Olympus Corp.) was then inserted via the working channel of the FB. The electrocautery snare was applied in a blended way at 12 W for cutting and coagulating the spherical part of the mass (Figure 2B). Because the mass base was too wide to be removed by the electric snare, it was ablated using a low-temperature plasma therapy device (EIC7070-01; Smith and Nephew, London, United Kingdom) under the laryngoscope (Figure 2C). To prevent a tracheoesophageal fistula, the base of the tumor was not completely removed (Figure 2D). The extracorporeal membrane oxygenation team was present to administer emergency treatment if intubation was difficult or asphyxia occurred.
Figure 1 Preoperative and postoperative imaging. A: Computed tomography (CT) revealed a mass occluding about 90% of the tracheal lumen; B: Contrast-enhanced CT a soft tissue mass with obvious enhancement on the right posterolateral wall of the trachea; C: CT indicated a slight protrusion at the position of the incision.

Figure 2 Fiberoptic bronchoscopy findings. A: A mass was located in the trachea at the 4- to 7-o’clock position about 1 cm below the vocal cords; B: The spherical part of the mass was removed by an electrocautery snare; C: The base was ablated by a low-temperature ion probe; D: Most of the base was removed; E: There was a slight protrusion at the position of the incision.

Pathological results
The tumor consisting of bundles of spindle cells with slender palisade nuclei, as shown by staining with hematoxylin and eosin (Figure 3A). Immunohistochemical analysis showed that the positive indicator of the tumor was S-100, and the results of CD117, desmin, smooth muscle actin, and CD34 were negative for the tumor (Figure 3B). The histological and immunohistochemical findings were consistent with benign schwannoma.
Figure 3 Pathological images of tumor cells. A: A histologic section from the tumor shows that the tumor cells were spindled with elongated palisading nuclei (hematoxylin–eosin stain, 400 ×); B: The tumor cells showed diffuse and positive S-100 staining by immunohistochemistry (streptavidin peroxidase method, 200 ×).

OUTCOME AND FOLLOW-UP

The patient’s clinical manifestations disappeared after removal of the tumor, and the tracheotomy cannula was successfully removed 1 wk after the operation. Repeat bronchoscopy and chest CT examinations were conducted for 6 mo after the procedure (Figure 1C). A slight protrusion was present at the position of the incision (Figure 2E).

DISCUSSION

Neurilemmomas, also known as schwannomas, originate from Schwann cells in the peripheral nerve tissues and are usually benign. They are more commonly reported in the head, retroperitoneum, neck, lungs, and extremities[2]. However, tracheal neurilemmomas are rare in children.

In general, neurilemmomas grow slowly. Because their clinical presentation is nonspecific and insidious, diagnosis is often delayed. Only when the tumor attains a large size can patients be diagnosed. The main symptoms include coughing, wheezing, and dyspnea[4]. Pulmonary function testing is an effective method to distinguish tracheal neurilemmomas from asthma[5,6]. Chest CT or magnetic resonance imaging can be used to measure the tumor’s size, location, and degree of intratracheal and extratracheal invasion. In this case, we reviewed the examination results before admission. Two months previously, we had discovered a mass in the upper part of the trachea. Lung function testing had also indicated an inspiratory phase obstruction, but it was initially ignored. In patients with an atypical presentation of asthma but with no response to anti-asthma treatment, bronchoscopy can detect airway tumors early and avoid misdiagnosis.

Because of the high degree of airway obstruction in this patient, extensive preoperative planning was necessary in case of an airway emergency during the operation. We planned to insert a 4.5-mm endotracheal tube under bronchoscopic guidance after anesthetic induction. Because the endotracheal intubation affected the exposure of the tumor, it would have been difficult to repeat the intubation in case of intraoperative decannulation. The 4.5-mm tube could not prevent blood from flowing into the lower airway, so we performed a tracheotomy below the tumor. If it had been difficult to intubate the patient by bronchoscopy, we would have attempted rigid bronchoscopy. Once a patient has developed obvious airway obstruction or asphyxia after induction of anesthesia, emergency tracheotomy or extracorporeal membrane oxygenation should be considered.

The tumor’s size, location, depth of tracheal wall invasion, and extratracheal involvement should be considered when treating primary tracheal neurilemmoma[7]. The main treatment strategies are surgical resection and endoscopic excision[8]. Traditionally, the best conduit for the management of endotracheal tumors is probably rigid bronchoscopy[8]. Because the tumor in our patient was in the upper trachea and occupied most of the airway, we initially attempted to use an electric snare to remove the tumor. Electrocautery snares can sever large masses quickly with concomitant hemostasis. However, because the base of the tumor was broad, the snare could not remove it. We chose coblation to remove the base of the tumor and reduce the probability of recurrence after electrocautery snaring. The advantages of coblation
include rapid and precise ablation, minor thermal damage, and the integrated function of suction and coagulation[9]. Because of these characteristics of coblation technology, there was minimal surrounding tissue damage, which minimized the likelihood of postoperative stenosis.

Although a small part of the tumor remained after its removal, the patient displayed no symptoms and her parents were opposed to further invasive open surgery. Nearly one-fourth of patients who undergo endoscopic resection develop recurrence. The time of recurrence is variable, but there is a chance of late recurrence. Given that these tumors grow slowly, it is preferable to perform yearly bronchoscopic observations. If the tumor continues to grow, tracheal resection is the best option. Outcomes and complications, such as airway stenosis and tumor recurrence, should be consistently evaluated under long-term follow-up.

CONCLUSION

Intratracheal neurilemmoma is relatively rare in children. Primary symptoms include coughing, wheezing, and dyspnea. Chest CT can be used to measure the tumor’s size, location, and degree of intratracheal and extratracheal invasiveness. The primary treatment strategies used for tracheal neurilemmoma are surgical resection and endoscopic excision. Long-term follow-up is warranted for the evaluation of outcomes and complications.

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