Persistent Tracheo-cutaneous Fistula: A Case Report

Persistan Trakeo-kutanöz Fistül: Olgu Sunumu

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Abstract
Persistent tracheo-cutaneous fistula, or persistent tracheal stoma, is a potential late complication of a tracheostomy. It commonly occurs in children due to a failure of spontaneous closure after decannulation. In adults, however, this is relatively less common. Described is a case of a 38-year-old man who, despite undergoing early and successful decannulation, presented with a persistent tracheal stoma due to pulmonary tuberculosis.

Key words: Tracheo-cutaneous fistula, persistent tracheal stoma, pulmonary tuberculosis, anti-tuberculosis treatment.

Tracheostomy is a procedure in which a surgical airway is created in the cervical trachea. It is commonly done in patients who have difficulty when removed from a ventilator, when there is an inability to protect the airway because of trauma or severe neurological damage, there is an upper airway obstruction that needs to be emergently bypassed, a life-threatening infection of the upper airway, or congenital upper airway anomalies (1).

Immediate complications can include apnea, bleeding, and pneumothorax/pneumomediastinum. Potential late complications include tracheoinnominate fistula, tracheomalacia, tracheal stenosis, tracheo-esophageal fistula, and tracheo-cutaneous fistula (TCF).

Decannulation is performed after successful tracheostomy tube plugging overnight. The wound heals and closes spontaneously in 5 to 7 days. When the tracheostomy tract undergoes epithelialization from the skin to the tracheal mucosa, a stoma may persist as TCF. It is a delayed complication of tracheostomy, especially in the pediatric age group, with an incidence of 3.3% to 43%, but is relatively less common in adults (2-5).

The most important factor that contributes to the persistence and failure of closure is the duration of cannulation (6,7). Another predisposing factor is the starplasty tracheostomy technique, which was introduced in 1990 as an alternative tracheostomy technique in children (8). A tracheo-cutaneous fistula...
communication is intentionally created by forming a circumferential muco-cutaneous suture line between the skin and the tracheal mucosa (6).

The other predisposing factors are poor nutritional status, immunosuppression (high-dose steroids, etc.), radiotherapy, infection, granulomatous disease, distal obstruction due to bilateral vocal cord paralysis, and tracheal stenosis (9,10).

Patients with TCF may suffer from recurrent aspiration and subsequent respiratory infection, difficulty in phonation, difficulty in clearing secretions, ineffective cough, difficulty swallowing, skin irritation and/or ulceration, cosmetic and social issues, and intolerance to swimming (11). This may lead to considerable discomfort, especially after successful decannulation.

Management of TCF is usually achieved with surgical excision of the fistula tract. Two techniques are available. Primary closure with drain placement offers better cosmetic healing of the scar with a higher risk of subcutaneous emphysema. Secondary closure involves excision of the fistula, replacement of a small tracheostomy tube, and subsequent decannulation, thereby ensuring healing by secondary intention and closure. This is not often done, as it is less acceptable to the patient to have another tracheostomy after decannulation, but it is a relatively safer procedure.

CASE

A 38-year-old male was attended to in the respiratory medicine department. He had undergone an emergency craniotomy and evacuation of an acute fronto-parieto-temporal subdural hematoma 2 months earlier. Due to difficulty with removal of the ventilator, a tracheostomy was performed. He was successfully decannulated after 3 weeks. He had a persistent opening and non-foul-smelling discharge from the tracheostomy site for 2 weeks. He was referred for evaluation of the trachea to rule out tracheomalacia and tracheal stenosis before a planned suturing of the TCF. There was also a history of a non-productive cough for 4 months prior to the surgery associated with weight loss and anorexia, but he had no fever and attributed his debilitation to the head injury he sustained a year earlier.

The patient had no history of tuberculosis, diabetes mellitus, hypertension, or other significant comorbidities. He was a non-smoker and only occasionally consumed alcohol. He was an electrician by occupation.

Clinical examination revealed a conscious, cooperative patient in no acute distress, with stable vital signs with a body mass index of 16.7 kg/m². He had a 2x2 cm anterior midline opening in the suprasternal area with non-foul smelling, purulent discharge at the tracheostomy site (Figure 1A). His lungs were clear on auscultation and other systems were within normal limits. His chest X-ray showed bilateral upper zone air-space opacities (Figure 2).

Figure 1A and B: Examination of the neck: A (top) Tracheostomy site, inflammation and purulent discharge; B (bottom) Tracheostomy, healed and closed, after 2 months of anti-tuberculosis therapy

Figure 2: Chest X-ray: Right upper zone, Left upper zone opacities

He had also received a 2-week course of antibiotics for methicillin-sensitive Staphylococcus aureus grown in pus
culture from the tracheal stoma. Since the tracheostomy wound had persisted and continued to emit discharge, and as he could not expectorate despite sputum induction, a bronchoscopy was done, which revealed unhealthy necrotic mucosa and purulent discharge around the fistulous opening and purulent secretions in the right and left upper lobe bronchi (Figure 3A). The secretions from the tracheal opening and from the upper lobes were separately sent for microbiological examination (Gram stain and cultures, acid-fast bacilli [AFB] stain, and cartridge-based nucleic acid amplification test [CB-NAAT]). Both samples showed numerous AFB, and Mycobacterium tuberculosis was detected in the CB-NAAT.

An anti-tuberculosis chemotherapy regimen of 2 (HREZ) followed by 4 (HR). After 6 months of antituberculosis therapy, the tracheal stoma closed (Figure 1B) with no tracheomalacia or stenosis (Figure 3B). His chest X-ray showed good radiological clearance and his sputum was negative for AFB.

**DISCUSSION**

While TCF is a well-known potential late complication of a tracheotomy in the pediatric age group, in adults it is relatively less common and easier to manage. The incidence rate of TCF from studies of children undergoing a tracheostomy varies from 3.3% to 43%. In the 1960s, the incidence of TCF was low, at just more than 3% (2,3). The incidence has increased with the increased number of indications for a tracheostomy (4,5).

According to Kubler and Passy (6), the incidence of persistent tracheal stoma is 70% if the cannulation period is 16 weeks or more, while tracheostomies close spontaneously when less than 16 weeks. In a retrospective study of 164 children performed by Ha et al. (12), the relative risk of TCF in a case of tracheostomy was significantly greater when longer than 24 months (risk ratio: 2.52) than for less than 12 months.

In a study conducted by Grønhøj et al. (13), neuromuscular disease was the most common indication. Of 69 tracheostomized children, 9 developed TCF or wound granulation and 53% of lower airway infections were due to Staphylococcus aureus. Nassif et al. (14) reported that of 57 patients who underwent a tracheostomy, 39 (68%), were for upper airway obstruction and 18 (32%) were due to prolonged ventilation. Nine cases (33% of decannulations) had TCF that required surgery. Al-Samri et al. (4) reported that upper airway obstruction due to subglottic stenosis and craniofacial syndromes were the 2 most common indications in 15 (21%) patients each, and TCF was seen in 37%.

Huber et al. (15) have also reported a case of a patient with burns whose TCF healed with local wound care and without surgery.

While granulomatous infection is a known predisposing factor, cases with pulmonary tuberculosis leading to failure of spontaneous closure of a tracheostomy have not been widely reported in the literature. However, Tong and Chow (16) reported a case of tuberculous tracheitis in the absence of pulmonary parenchymal lesion presenting as TCF, which resolved with anti-tuberculosis treatment.

Our case is an example of a chronic, indolent infection, tuberculosis, as a cause of the lack of a spontaneous closure of the tracheostomy. In developing countries, tuberculosis constitutes a major healthcare burden with significant morbidity and mortality. Complications of surgical sites and tracheostomies by tuberculosis must therefore be considered in any case of delayed wound healing. Early detection and treatment can significantly
reduce morbidity and mortality and avoid surgical closure in some cases.

CONCLUSION
Our case demonstrates an important cause of the failure or delayed closure of a tracheostomy after decannulation – pulmonary tuberculosis. The tracheostomy site bears the brunt of contaminants from the oral cavity and the lungs. The likely mechanism for tracheal stoma infection is retrograde spread of M. tuberculosis bacilli from the pulmonary focus of infection, especially when the duration of cannulation is short, cannot be over-emphasized. When treated, it may obviate the need for surgery. Our case also represents the tip of the iceberg of patients with undiagnosed pulmonary tuberculosis who are a public health challenge, as they come in contact with family, healthcare providers, and society at large. In our case, universal health precautions were followed, as always. Additionally, if a patient with pulmonary tuberculosis requires life-saving emergency surgery, the operation theater will be subjected to sterilization/disinfection procedures.

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CONFLICTS OF INTEREST
None declared.

AUTHOR CONTRIBUTIONS

REFERENCES