Dysphagia in a Young Adult: Rare Case of Giant Cervical Osteophyte

Abstract
Cervical osteophytes may be seen in diffuse idiopathic skeletal hyperostosis, ankylosing spondylitis, posttraumatic, postoperative, degenerative causes, cervical spondylosis, and infectious spondylitis. A cervical osteophyte is very rarely considered among the differentials for symptoms of dysphagia. C5–C6 as well as C6–C7 being a site of greater load-bearing and mobility, the propensity to form osteophytes is high, with a small osteophyte leading to local mass effect. A 42-year-old male patient presented with mild dyspnea and significant dysphagia since 8 months, accompanied by dysphonia, weight loss, and intermittent aspiration. Clinical examination including neurological examination was normal. A barium swallow showed that osteophytes were severely protruding and displacing the lower pharynx and the proximal esophagus anterosuperiortly. The patient underwent surgical removal of the osteophyte through Smith–Robinson approach. Complaints of dysphagia were significantly decreased in postoperative period. A thorough evaluation is necessary to rule out other causes of dysphagia. Surgical management of this uncommon condition might be considered after confirmation of the osteophyte to be the offending lesion as it has favorable clinical outcomes.

Keywords: Dysphagia, giant cervical osteophyte, management

Introduction
Ossification and calcification of soft tissue and formation of osteophytes involving predominantly the anterior longitudinal ligament[1] is seen in about 30% of adults over the age of 60 years, with the incidence increasing with age.[2] Cervical osteophytes may be seen in diffuse idiopathic skeletal hyperostosis, ankylosing spondylitis, posttraumatic, postoperative, degenerative causes, cervical spondylosis, and infectious spondylitis.[3] The symptoms usually relate to surrounding structures when compressed. Dysphagia is the most common symptom and rarely nonspecific pain, decreased neck range of motion, stridor, dyspnea, dysphonia, hoarseness, foreign body sensation on deglutition as well as sleep apnea.[4,5] A cervical osteophyte is very rarely considered among the differentials. Hence, they are typically investigated after other factors causing dysphagia are ruled out. C5–C6 as well as C6–C7 being a site of greater load-bearing and mobility, the propensity to form osteophytes is high.[6] A large osteophyte at C4–C7 levels can lead to substantial mechanical constriction of esophagus. A small osteophyte coupled with inflammation at cricoid level may lead to local mass effect.[7] Usually managed conservatively, for patients refractory to conservative management or with increased severity of symptoms operative management is advised. Cervical osteophyte causing dysphagia is generally seen in the elderly population. We report a case of a young male who presented with complaints of increasing dysphagia as well as change in voice and was diagnosed with giant anterior cervical osteophyte after a barium swallow by a gastroenterologist.

Case Report
A 42-year-old male patient presented with mild dyspnea and significant dysphagia since 8 months. Gentle exertion and daily routine activities lead to shortness of breath. These symptoms had become worse around the time of his visit and were accompanied by dysphonia, weight loss, and intermittent aspiration. He complained of foreign body sensation. Various physicians failed to make an accurate diagnosis and the symptoms of the patient continued to deteriorate progressively.

The patient had no significant past surgical or medical history other than being a
smoker for the past 20 years. There was no abnormality of the tongue and oral cavity including hypopharynx on examination by an orthonathologist. Clinical examination including neurological examination was normal with no evidence of dysarthria.

No intracranial pathology was seen on magnetic resonance imaging (MRI) brain, done to rule out intracranial pathology. A barium swallow showed that osteophytes were severely protruding at the anterior portion of the 4th–6th cervical spine and displaced the lower pharynx and the proximal esophagus anterosuperiorly. In the oral phase, there was no abnormality other than premature bolus loss. In the pharyngeal phase, esophagus was displaced anterosuperiorly with constriction anterior to C5 vertebra. X-ray and computerized tomography (CT) scan of the cervical spine were done to confirm anterior cervical osteophytes and revealed the formation of large osteophytes anteriorly from 4th to 6th cervical vertebrae [Figures 1 and 2]. Osteophyte of the 5th cervical spine was the most prominent. It protruded 11 mm beyond the vertebral border. C-spine MRI was consistent with anterior herniation of the intervertebral disc with ruptured anterior longitudinal ligament. A T2-weighted image revealed high signal intensity with irregular enhancement. This could be a result of retropharyngeal irritation and inflammatory changes.

The patient had developed severe anxiety and sleep disturbances probably due to increasing severity of symptoms and lack of diagnosis. The patient underwent surgical removal of the osteophyte through Smith–Robinson approach. It included removal of the anterior osteophytes, C4–C5, C5–C6 cervical discectomy, interbody stand-alone polyetheretherketone cage as well as bone graft. Complaints of dysphagia were significantly decreased in postoperative period [Figures 3 and 4]. He was started on normal diet 1 day after surgery. He also noticed a change in voice back to normal by 7 days. On further follow-up of 1-year, the patient was symptom free.

**Discussion**

Dysphagia associated with cervical anterior osteophytes occurs mainly due to mechanical compression and displacement of surrounding structures. Concomitant pharyngoesophageal inflammation and fibrosis may further aggravate the situation. Dysphagia is often seen with solid foods and is usually progressive. Conservative treatment methods are encouraged primarily. Patients refractory to conservative modalities are treated surgically with excision of osteophyte with or without fusion of the affected segments.

Huge osteophytes located at the upper cervical spine may at times cause difficulties in intubation by narrowing the hypopharynx as well as displacement of the epiglottis and cords whereas osteophytes located at the lower levels may do so by pushing the trachea and larynx forward. Anesthetists must be mindful of this, and fiberoptic or videolaryngoscopic intubation should be preferred. Tracheostomy may be rarely needed due to possible airway problems preoperatively and postoperatively.

In the present case, besides the direct bony compression, the inflammation and edema in the retro-laryngopharyngeal space may also have been a cause for acute exacerbation of symptoms. The diagnosis and initiation of appropriate treatment was delayed probably since osteophyte induced dysphagia is very rare in young individuals. Simultaneous mild exertional dyspnea and dysphonia as well as absence of neck pain or cervical radiculopathy compounded this. If there is a presence of respiratory symptoms with dysphonia, neurological causes should be ruled out through, adequate investigations as was done for our case.
Giant cervical osteophytes leading to dysphagia is not uncommon. It must be considered in elderly as differential diagnosis of dysphagia. CT scan is required to confirm diagnosis, evaluate, and clearly define the bony anatomy. A barium swallow is essential to confirm the presence of esophageal compression by the cervical osteophyte and was the primary investigation which leads to diagnosis in our case. The presence of dyspnea and dysphonia, calls for MRI. MRI is also helpful to see coexisting spinal cord or root compression. Some authors opine that osteophytes causing chronic dysphagia and dyspnea must be excised, because of possible progression to acute respiratory distress. Surgical excision has good long-term outcomes. We performed fusion at both the levels as there was rupture of the anterior annulus of the disc and concomitant disc degeneration. Fusion may regress the formation of cervical osteophyte by taking care of abnormal mobility at the site. Our patient is symptom free at 1-year follow-up with minimal radiographic regrowth [Figures 5 and 6].

**Conclusion**

A thorough evaluation is necessary to rule out other causes of dysphagia. Surgical management of this uncommon condition might be considered after confirmation of the osteophyte to be the offending lesion as it has favorable clinical outcomes.

**Declaration of patient consent**

The authors certify that they have obtained all appropriate patient consent forms. In the form the patient(s) has/have given his/her/their consent for his/her/their images and other clinical information to be reported in the journal. The patients understand that their names and initials will not be published and due efforts will be made to conceal their identity, but anonymity cannot be guaranteed.

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Conflicts of interest

There are no conflicts of interest.

References