Nonsurgical treatment of stylohyoid (Eagle) syndrome: a case report

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Abstract (J Korean Assoc Oral Maxillofac Surg 2014;40:246-249)

Eagle syndrome is a rare condition caused by elongation of the styloid process or calcification of the stylohyoid ligament. Patients with Eagle syndrome typically present with dysphagia, dysphonia, cough, voice changes, otalgia, sore throat, facial pain, foreign body sensation, headache, vertigo, and neck pain. Here we report a case in which the patient initially presented with sore throat, left-sided facial pain, and cough. This case report provides a brief review of the diagnosis and nonsurgical management of this rare syndrome.

Key words: Eagle syndrome, Dysphonia, Heterotopic ossification, Temporal bone

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I. Introduction

Eagle syndrome is a rare clinical condition caused by elongation of the styloid process or by mineralization (ossification or calcification) of the stylohyoid ligament. The length of the styloid process is normally 2 to 2.5 cm, and elongation beyond 2.5 cm causes Eagle syndrome, which was first defined in 1937.

This syndrome has two types, including classic and styloid-carotid syndromes. The classic syndrome, also known as stylalgia, is usually characterized by pharyngeal pain localized to the tonsillar fossa, referred otalgia, and neck pain. It may also be associated with dysphagia, hypersalivation, sensation of a foreign body in the throat, and transient voice changes that are often seen following tonsillectomy. Styloid-carotid syndrome is characterized by nonspecific symptoms caused by compression of the carotid arteries and sympathetic fibers by the styloid process. The most common etiology of this syndrome is ossification or calcification of the stylohyoid ligament.

II. Case Report

A 53-year-old woman was referred to the pain clinic at Amir Alam Hospital with a complaint of pain in the throat and the left side of the head and face. Her pain began 16 years prior following a root canal of a mandibular tooth on the left side. She at times experienced severe pain attacks lasting approximately 10 minutes with a frequency of 10 times per day, which extended from the external auditory canal to the left side of the throat. Additionally, cough was a trigger for these pain attacks. Certain odors or nervousness were also aggravating factors of her pain, while pressure on the ear and diphenhydramine were mitigating factors. The patient reported her highest and lowest pain intensity as 4 and 2, respectively, according to the numeric rating scale (NRS). The patient had a past medical history significant for hypertension and favism.

The patient’s general health as well as her head and neck
were thoroughly examined. Her skull, cranial nerves, eyes, ears, nose, sinuses, thyroid, and dentition were unremarkable. No cervical, axillary, epitrochlear, or inguinal adenopathy was detected. On oral and pharyngeal examination, severe tenderness was present on a bony prominence in the tonsillar cavity. Head and neck rotation was painful at the end of active range of motion. As a result, lateral skull and neck x-rays were taken that showed elongation of the styloid process, thereby confirming the diagnosis of Eagle syndrome. (Fig. 1) Laboratory studies and histopathology were not performed.

Upon diagnosis, treatment with pregabalin (75 mg/day) and amitriptyline (10 mg/day) was initiated. These medications improved the patient’s cough and reduced the pain severity by about 50% within 2 months. Three and 6 months after treatment, her pain severity improved by about 80% and her NRS score decreased to one. The patient also experienced occasional pain-free intervals. (Table 1)

### III. Discussion

In this study, Eagle syndrome was identified in our patient, after which medical therapy was initiated with an appropriate response.

Eagle syndrome may be unilateral or bilateral, although in most cases symptoms are unilateral. According to Moon et al.10, Eagle syndrome and elongation of the stylohyoid ligament is typically a bilateral process, though in our patient this condition was unilateral.

A calcified stylohyoid ligament is found in 4% to 28% of normal population11. There is no direct correlation between the severity of Eagle syndrome and level of calcification12. However, there is a clear relationship between longer lengths of the styloid process and higher pain intensity and severity of Eagle syndrome8.

Pharyngeal pain with radiation to the neck and ears poses a very difficult challenge for interpretation, including a vast number of differential diagnoses13-17. (Table 2) The differential diagnosis for Eagle syndrome includes inflammatory disorders, masses of the pharynx and tongue base, and cranial nerve neuralgia. On the other hand, patients with an elongated styloid process manifest some transient and nonspecific symptoms as well as the severe classic symptoms, which require surgery18. Traumatic fracture of the apophysis, pharyngeal infection and inflammation, rheumatoid disorders of the hyoid bone, inflammation of the muscles attached to the styloid process, osteoarthritis of the cervical vertebrae, pressure on adjacent nerves such as cranial nerve IX and the

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**Table 1. Patient history, work-up, treatment, and outcomes**

| Work-up            | Steps                                | Patient data                                                                 |
|--------------------|--------------------------------------|-------------------------------------------------------------------------------|
| History            | Age, gender, occupation              | Fifty-three years, female, housewife                                          |
|                    | Pain location                         | Throat and left side of head and face                                         |
|                    | Quality and severity                 | Severe and penetrating pain with a maximum numeric rating scale of 4          |
|                    | Alleviating factors                  | Pressure on ear and diphenhydramine                                          |
|                    | Aggravating factors                  | Certain smells and nervousness                                                |
|                    | Setting and radiation                | Coughing fits are associated with her pain attacks                            |
|                    | Past medical history                 | Hypertension and favism                                                       |
| Family history     | Medications                           | None                                                                          |
| Physical examination| Head, neck, eyes, ears, nose, throat | Diphenhydramine and ibuprofen                                                 |
| Imaging            | Lateral skull and neck X-ray         | Severe tenderness on a bony prominence in the tonsillar cavity                |
| Treatment          | Medical treatment                    | Elongation of the styloid process                                             |
| Outcome            |                                      | Pain improvement and numerical rating scale: 1                               |

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mandibular branch of the 5th cranial nerve and corda tympani, pathologies of the third molar, and pharyngeal scar following tonsillectomy are other pathologies that could mimic symptoms of Eagle syndrome. Different forms of pharyngeal neuralgia may also result in similar symptoms including laryngeal neuralgia, occipital neuralgia (involving Arnold’s nerve), sphenopalatine neuralgia (secondary to sphenoiditis inflammation), and finally trigeminal neuralgia which may account for sporadic pains with pressure within the auditory canal. Disorders of the temporomandibular joint constitute another possible diagnosis. In our case, most other differential diagnoses were ruled out according to the history and physical examination findings.

Clinical diagnosis rests upon previous history of trauma or tonsillectomy and palpation of the tonsillar fossa. Radiologic studies such as an orthopantomograph or lateral skull view with the head slightly extended may help to confirm this diagnosis. A precise history, examination, and imaging studies also contribute greatly to achieving the correct diagnosis. (Table 2) The final diagnosis in our case was confirmed by imaging.

Medical therapy is first-line treatment for Eagle syndrome. Surgery with resection of the elongated styloid process is considered to be definitive treatment, however, surgery may be contraindicated in some cases or patients may decline operative intervention. In these cases, conservative treatment with oral medications may be considered. In cases that do not respond to medical therapies, surgery is indicated. Different medications may be used in medical management of Eagle syndrome based on the respective etiology, including analgesics, anticonvulsants, antidepressants, and local infiltration with steroids or long-acting local anesthetic agents. In our case, a tricyclic antidepressant (amitriptyline) and an anticonvulsant (pregabalin) were started after diagnosis. Our patient dramatically and persistently responded to conservative treatment after 3 and 6 months of therapy. Nonsurgical treatment of Eagle syndrome with gabapentin, tianeptine, tramadol, acetaminophen, local lidocaine injection and stellate ganglion block has also been reported. However, to our knowledge, our patient is the first case of Eagle syndrome to be treated with only oral medications, while previous similar cases also utilized a stellate ganglion block.

The exact mechanism by which medications achieve symptom relief in patients with Eagle syndrome is not known.

Table 2. Differential diagnoses for pain in the head, cervicofacial, and cervicopharyngeal regions

| Etiology                          | Differential diagnosis                                                                 |
|----------------------------------|----------------------------------------------------------------------------------------|
| Vascular3                        | Migraine, cluster headache, chronic tension and cervicogenic headaches, carotidynia, atypical facial pain, paroxysmal hemierciasia; headaches of reactive vasodilation: fever, drug-induced, postictal, hyperthyroidism, hypoglycemia, hypoxia, hypercarbia; headaches associated with arterial hypertension: chronic severe hypertension, pheochromocytoma, coital headaches; headaches caused by cranial arteritis: temporal arteritis, etc. |
| Muscle spasm1,14                  | Headache of posturally-induced or perilesional muscle spasm: impaired posture, cervical spondylosis and other diseases of cervical spine; myofascial pain dysfunction syndrome (headache or facial pain associated with disorders of teeth, jaw, and related structures, or TMJ syndrome); headaches associated with psychophysiological muscular contraction: muscle contraction headaches or tension-type headaches associated with disorder of periracinal muscles |
| Without demonstrable physical substrate | Heads of uncertain etiology: tension headaches unassociated with disorder of periracinal muscles, some forms of posttraumatic headache; psychogenic headaches: hypochondriacal, conversional, delusional, and malingered; facial pain of uncertain etiology: atypical facial pain |
| Combined tension-migraine        | Episodic migraine superimposed on chronic tension headaches, chronic daily headaches (associated with analgesic and/or ergotamine overdose, also called rebound headaches; not associated with drug overuse) |
| Meningeal inflammation           | Subarachnoid hemorrhage, meningitis and meningoencephalitis, meningeval carcinomatosis |
| Altered intracranial pressure    | Increased intracranial pressure: intracranial mass lesions (neoplasm, hematoma, abscess, etc.), hydrocephalus, benign intracranial hypertension, venous sinus thrombosis |
| Cranial neuralgia1,16            | Decreased intracranial pressure: postlumbar puncture headaches, spontaneous hypoliquorheic headaches |
| Bones and joints1,14              | Postherpetic neuralgia, glossopharyngeal, trigeminal, superior laryngeal, occipital, pterygopalatine ganglion, intermediate nerve, geniculate neuralgia |
| Ear, nose, and throat diseases    | Chronic tonsillitis, tonsillar calculus, spasm of the pharyngeal constrictor muscle, otitis, mastoiditis, fracture of the hyoid bone, pterygoidhamulus burstsis |
| Other diseases17                  | Chronic laryngopharyngeal reflux, psychosomatic diseases, foreign bodies, inflammatory and neoplastic processes in the oropharyngeal area, pharyngeal and base of tongue tumors, nuchal cellulitis and fibrosis, neck-tongue syndrome |
| Referred pain                    | TMJ pain, cardiac pain, diaphragmatic irritation, gastrointestinal sources (peptic ulcer disease, gallbladder, pancreas) |

(TMJ: temporomandibular joint)

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The cause of pain in Eagle syndrome is the stimulation of adjacent nerves by the elongated styloid process and secondary induced inflammation. It seems that medications such as anticonvulsants and antidepressants may reduce nerve stimulation and consequently pain intensity by altering the concentration of neurotransmitters, and analgesics such as nonsteroidal antiinflammatory drugs may improve pain by reducing inflammation in adjacent tissues.

In conclusion, lateral skull imaging in cases suspicious for Eagle syndrome is recommended to confirm this diagnosis, and medical therapy should be considered as first-line treatment for this rare condition.

Conflict of Interest

No potential conflict of interest relevant to this article was reported.

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