Eagle’s syndrome: an unusual cause of recurrent neck pain in a child

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DESCRIPTION
A 14-year-old girl presented with recurrent episodes of pain over the right side of the neck for the past 1 year. It was located in the upper neck, dull aching, moderate in intensity (6–7/10 on the pain scale) and radiating to the ipsilateral lower jaw, ear and face. There were 2–3 episodes/day and each episode lasted for 10–15 min. It occasionally aggravates on taking solid food and turning the head towards the left side. She underwent bilateral tonsillectomy one and a half months back and her symptoms had aggravated following the procedure. On examination of the head and neck, no external abnormality was seen. Palpation elicited pain over the right submandibular area anterior to the sternocleidomastoid muscle and intraorally in right retromolar trigone region and peritonsillar fossa. X-ray (figure 1) and CT head (figure 2) showed a bilaterally elongated styloid process, confirming the diagnosis of Eagle’s syndrome (ES). Child was planned for the surgery, but parents were not ready for the procedure. She started on oral amitriptyline (25 mg/day) and intermittent analgesics. At 3 months follow-up, she had partial improvement in her symptoms (both intensity and frequency of the episodes was decreased).

ES is a rare condition secondary to elongated ossified styloid process and was described by Eagle in 1937. Females of third to fourth decades are more commonly affected and only five patients have been reported in children so far. The accepted length of the styloid process is 2.5 cm and greater length is considered abnormal. While styloid abnormalities often occur bilaterally but pain symptoms usually present unilaterally. Pain is the most common presenting symptom of ES and is often unilateral, on the side of the elongated styloid process. Sometimes it radiates to the ipsilateral ear and jaw and present as otalgia or temporomandibular joint pain and may aggravate with yawning and head-turning. Other symptoms are dysphagia/odynophagia, voice changes and foreign body sensation. A patient may experience visual disturbance, syncope, ischaemic stroke and rarely Horner syndrome due to carotid artery compression. Presence of an elongated styloid process alone or the pain syndrome alone, are not consistent with the diagnosis of ES and both an elongated styloid process and an appropriate clinical picture is required.

Eagle divided ES into two subtypes: classical ES and stylocarotid syndrome. Classical ES presents with pain, dysphagia and a foreign body sensation that presents immediately after tonsillectomy while pain, visual disturbance and syncope due to carotid artery compression are the features of stylocarotid syndrome. However, further studies showed that it can occur without tonsillectomy.

The aetiology and pathogenesis of ES are still controversial. Trauma to styloid process, retained embryological cartilage tissue from Reichert’s cartilage, calcification of the stylomandibular ligament and expansion of osseous tissue at the origin of the stylomandibular ligament are the proposed theories. Neuropathy of sphenopalatine ganglia, submandibular sialolithiasis/sialadenitis, glossopharyngeal and trigeminal neuralgia, chronic tonsillitis, pain due to tonsillectomy while pain, visual disturbance and syncope due to carotid artery compression are the features of stylocarotid syndrome. However, further studies showed that it can occur without tonsillectomy.

Conservative management of ES includes analgesics and a combination of anticonvulsants, antidepressants, local anaesthetic/steroid injections. Literature tends to support surgical treatment as a more definite treatment with long-lasting symptomatic relief. Surgical treatment includes resection of the elongated styloid process via an intraoral or cervical approach. To conclude, ES is very rare in the paediatric population, but it
should be kept in the differential diagnosis of recurrent neck and cranial pain.

Learning points

► Eagle’s syndrome (ES) is a rare condition secondary to elongated ossified styloid process and neck pain is the most common presenting symptom.
► It is a rare case of recurrent neck pain in children and only five patients have been reported so far.
► Glossopharyngeal and trigeminal neuralgia, submandibular sialolithiasis, sialadenitis, hyoid bursitis, otitis media, temporal arteritis and temporomandibular arthritis are the differentials of ES.

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REFERENCES