Clinical Reasoning: Forty-Year-Old Woman With Scapular Winging and Dysphonia

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M. Aladawi: contributed with Drafting and revision of the manuscript, including medical writing for content, acquisition of data, study concept and design, and Analysis and interpretation of data
M. Punsoni: Contributed in medical writing for content, major role in the acquisition of data, and analysis and interpretation of data
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Section 1:

A 40-year old woman presented to the neuromuscular clinic for evaluation of chronic right shoulder weakness of 1-year duration. During that time, she had noticed progressive difficulties lifting her right arm overhead and reaching for objects on the back seat of her car. Around 3 months after the onset of shoulder weakness she started to develop an intermittent achy sensation behind the right ear and hoarseness of her voice. She noted difficulty talking over loud noises and in phone conversations. She denied vision, taste, or hearing changes, dysphagia, dyspnea, numbness, or weakness in other limbs. She did not have any prior shoulder trauma or past surgical history involving her neck. She denied severe pain in the shoulder region prior to the onset of symptoms. There was no family history of shoulder muscle weakness or gait impairment.

Neurological examination demonstrated normal mental status examination; patient speech was fluent, and she was able to name, repeat and comprehend. Of note, her voice was hoarse with a reduced pitch but all words were intelligible, and no nasal or strained quality was present. Cranial nerve examination showed reduced soft palate elevation on the right side. Gag reflex was present. Motor exam demonstrated right shoulder drop, atrophy of right trapezius muscle, lateral displacement of the right scapula at rest that worsened with shoulder abduction. Full active forward flexion of the right shoulder was limited to 120 degrees but improved to 180 when lying supine. Examination of individual muscle groups demonstrated 4/5 weakness in left lateral neck rotation, 5/5 neck flexion, and 4/5 weakness in right shoulder shrug. Sensory and deep tendon reflexes examination disclosed no deficits.

Questions:

Where would you localize this neurological process?

Section 2:

Neurogenic scapular winging can be due to long thoracic neuropathy, spinal accessory neuropathy or less commonly dorsal scapular nerve involvement. Importantly, neurological exam can point to the nerve involved. Serratus anterior muscle weakness (long thoracic nerve) causes medial winging of the scapula, which is more prominent with forward shoulder flexion. Scapular winging is lateral and worsens with shoulder abduction in lesions of spinal accessory nerve. Winging from rhomboid weakness (dorsal scapular nerve) presents with pain at the medial border of the scapula and becomes more prominent by having the patient push the elbow backward against resistance with the hand on the waist.

The combination of shoulder drop, trapezius atrophy, lateral scapular winging that worsens with abduction and weakness of ipsilateral sternocleidomastoid (neck flexion and contralateral neck rotation) suggested a spinal accessory neuropathy in our patient. Neurological examination also demonstrated voice changes consistent with dysphonia, which together with the finding of reduced palatal elevation, could point to vagus nerve dysfunction. Retroauricular pain could be suggestive of glossopharyngeal nerve dysfunction (innervates retroauricular skin) or non-specific referred pain.
To evaluate further her right scapular winging, patient was referred for EMG testing. In this study, standard motor and sensory nerve conduction studies of the upper limb were normal. Needle exam showed fibrillating potentials and reinnervation (large, long motor units with reduced recruitment) in the right trapezius muscle. Serratus anterior, rhomboids, infraspinatus, cervical paraspinal, and limb muscles were normal. Patient was also referred to ENT service for further evaluation of her dysphonia. Laryngoscopy revealed right true vocal fold paralysis.

Questions:

What etiologies are you considering?
What ancillary testing is indicated next?

Section 3:

The EMG results pointed to a right spinal accessory neuropathy or C3-C4 radiculopathy. Preserved sensory nerve action potentials, as in this case, normally suggest an abnormality proximal to the dorsal root ganglion. However, the spinal accessory nerve lacks a sensory branch to evaluate in nerve conduction studies so in the absence of cervical paraspinal muscle involvement the distinction between pre- and postganglionic pathology cannot be made. MRI C-spine was obtained, which did not reveal significant neuroforaminal stenosis in the upper cervical segments. At this point, the clinical picture was consistent with multiple cranial neuropathies. We suspected the cause of the scapular winging was a spinal accessory neuropathy but the etiology remained unclear.

Lesions of the spinal accessory nerve are most commonly iatrogenic (i.e. posterior cervical lymph node biopsy) and affect the distal portion of the nerve innervating trapezius but sparing sternocleidomastoid muscle. Proximal and intracranial lesions of the accessory nerve affect the sternocleidomastoid and trapezius muscles as was seen in this case.

MRI brain and neck-soft tissue was then obtained, which revealed an enhancing mass on the right jugular foramen suspicious for meningioma or schwannoma (Figure 1). Atrophy of the right trapezius and sternocleidomastoid muscles was also noted on imaging.

Patient underwent craniotomy and a tumor originating in the right glossopharyngeal nerve was resected. This tumor was compressing but not infiltrating the vagal and the spinal accessory nerves. Pathology revealed a benign nerve sheath tumor with mixed features of schwannoma and neurofibroma (Figure 2). Patient had rapid improvement of her voice in the next 3 months and slower but steady improvement of her scapular winging over the next year. Dynamic muscle transfer, which is typically performed in cases resistant to conservative treatment, was not required. For trapezius muscle palsy, this surgery involves the lateral transfer of the rhomboid major, rhomboid minor and levator scapulae tendons to replace lower, middle, and upper trapezius muscle fibers respectively. This improves the function of the trapezius muscle and stabilizes the scapula.
Discussion:

The case presented was due to a neoplastic spinal accessory neuropathy in the context of multiple cranial nerve deficits including dysphonia (vagus nerve) and retroauricular pain (glossopharyngeal nerve). This constellation is called Jugular foramen syndrome (Vernet syndrome)\(^4\). Our case emphasizes the fact that proximal spinal accessory neuropathy is typically not iatrogenic and compressive or tumoral lesions should be sought\(^5\).

The jugular foramen is split by a fibrous septum into two parts: a smaller anteromedial portion (pars nervosa) and a larger posterolateral portion (pars vascularis). The pars nervosa carries the inferior petrosal sinus, and the glossopharyngeal, vagus, and accessory nerves, while the pars vascularis contains the sigmoid–jugular complex. The hypoglossal canal lies posteromedially and in close proximity. Both anatomical structures lie within the intercondylar space\(^4\). Lesion confined to the jugular foramen can result in glossopharyngeal, vagus, and accessory nerve palsies. The diagnosis of neoplastic jugular foramen syndrome can be difficult because cranial nerve involvement develops gradually, in both a variable chronological fashion and variable impairment of each cranial nerve, as was the case in the patient presented.

Lesions affecting the jugular foramen can also extend beyond this structure to involve the hypoglossal canal. Collet-Sicard syndrome is a neurologic disorder that manifests with glossopharyngeal, vagus, accessory, and hypoglossal nerve palsies due to lesions affecting the jugular foramen and intercondylar space. Villaret syndrome is when patient presents with Horner’s syndrome in addition to glossopharyngeal, vagus, accessory, and hypoglossal nerve palsies, which signifies carotid sheath involvement\(^6\). Etiologies of Jugular foramen syndrome include primary tumors (meningioma, schwannoma, cholesteatoma), inflammation (meningitis, malignant otitis media), bone metastasis, carcinomatous meningitis, and internal jugular vein thrombosis\(^7,8\).

The pathology in our case revealed findings consistent with a “hybrid” benign nerve sheath tumor. The presence of capsule, axons situated peripherally within the lesion, and diffuse S100 immunopositivity all support schwannoma while the prominent collagen within the tumor and lack of prominent Verocay bodies suggested possible neurofibroma histology\(^9\).

Neurofibromas typically develop in skin as nodular or polypoid lesions\(^9\). Schwannomas most commonly affect the vestibulocochlear nerve followed by the trigeminal nerve\(^10\). Schwannomas can be distinguished from other primary intracranial tumors by certain radiological features. In this case, the jugular foramen mass demonstrated a dumbbell shape and prominent diffuse contrast enhancement, which are typical nerve sheath tumor radiological features\(^11\). Meningioma, on the other hand, typically presents a dural-tail sign on imaging\(^12\). The benign nature of the encountered mixed nerve sheath tumor, which seemed to cause deficits by compression rather than invasion or infiltration, explains the good recovery our patient had post-resection.

Parsonage–Turner syndrome (or neuralgic amyotrophy) is a common non-iatrogenic cause of scapular winging. In this syndrome, trapezius muscle is involved in 20% of the cases and can sometimes present
with associated dysphonia mimicking a partial Vernet syndrome\textsuperscript{13,14}. The muscle weakness in this inflammatory monophasic condition is typically preceded by severe shoulder pain and most commonly causes a more diffuse and patchy motor involvement of the long thoracic, suprascapular, axillary, and anterior interosseous nerves\textsuperscript{15}.

Scapular winging is a frequent referral to neuromuscular clinics and has a broad differential diagnosis. Detailed clinical exam and electromyography can be very informative. A neoplastic cause, although rare, has to be considered in unilateral progressive scapular winging and especially if no history of trauma or prior medical procedures is present.

**Appendix 1 Authors**

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<tr>
<th>Name</th>
<th>Location</th>
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References:


Figure 1: MRI examination of the brain

Legend: Coronal (A) and axial (B) T1W post contrast sequences with fat saturation. Right jugular foramen mass (1.7 cm x 1.6 cm x 2.3 cm) demonstrating typical dumbbell shape and prominent homogenous contrast enhancement (arrows).

![Figure 1](image1.png)

Figure 2: Pathology analysis of the jugular foramen tumor

Legend: H&E at 100x (A) shows a neoplasm with characteristic spindle cells and a collagenous stroma. Immunostaining at 100x demonstrates diffuse positivity for S100 (B) while CD34 (C) shows limited reactivity in tumor cells.

![Figure 2](image2.png)
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