CASE REPORT

Accessory flexor carpi ulnaris: case report and review of the literature

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INTRODUCTION

Most of the accessory muscles of the forearm described in the radiology literature are located either in the radial aspect of the forearm or towards the hypothenar eminence. We present an unusual case of an ulnar-sided distal forearm accessory flexor carpi ulnaris muscle presenting as a “pseudotumour” demonstrated with both ultrasound and MRI, rarely reported in the current surgical and anatomical literature. Given the location and relation to the ulnar nerve towards Guyon’s canal, the accessory muscle may also predispose to distal ulnar nerve entrapment.

CASE REPORT; CLINICAL PRESENTATION AND IMAGING FINDINGS

A 13-year-old girl was referred to our tertiary orthopaedic oncology centre with an ulnar-sided swelling at the volar aspect of the distal right forearm. This was more prominent on flexion of the wrist. Other than the swelling, there were no other clinical symptoms, and in particular no symptoms of ulnar nerve compression. On clinical examination, the “pseudotumour” was smooth and painless on palpation and measured approximately 5 cm in craniocaudal (CC) dimension. (Figure 1) It was located towards the distal forearm on at the ulnar side with prominence particularly elicited during wrist flexion with ulnar deviation. The patient was right hand dominant and a keen rock climber. There was no significant family history of note.

The patient underwent dynamic ultrasound that identified a well-defined hypoechoic structure of identical echotexture to adjacent muscle bellies at the site of clinical concern, just deep to a normal appearing and separate FCU. (Figure 2) It measured in the region of 5 × 1.3 × 0.5 cm in size (CC x transverse x anteroposterior). The ulnar nerve was demonstrated at its deep margin. MRI was undertaken, confirming the ultrasound findings (Figure 3A, B) with a small distal tendon inserting at the pisiform separate to and just radial to the larger FCU tendon insertion (Figure 3C, D). The muscle measured up to 7 cm in CC dimension with its origin at the ulnar aspect of the distal ulna, deep and anterior to the proximal extensor carpi ulnaris (ECU) muscle belly. No intrinsic muscle signal abnormality was demonstrated. The ulnar nerve was demonstrated with normal dimensions and signal intensity, deep to the distal aspect of the accessory muscle belly and proximal to Guyon’s canal.

DISCUSSION

There are several reports in the literature with regards anomalies of FCU with wide variance in their features. The presence of a supernumerary muscle belly is rare when compared to descriptions of anomalous insertions or small extra tendinous slips. Furthermore, there are inconsistencies in the literature with regard to the nomenclature of the...
supernumerary muscles that have included digastric FCU, accessory FCU, and flexor carpi ulnaris palmaris (FCUP). Bhardwaj et al recently described a new classification system for supernumerary FCU in the surgical literature in an attempt to consolidate and clearly classify the anomaly, so as the reporting of their incidence and clinical features could be accurately assessed. The classification was based both on the clinical appearances and probable embryological basis of the anomaly. They separated the different types of anomalies of FCU into three major categories that included; Type I—split tendons, where there is a single muscle with two tendons, Type II—digastric FCU, where two heads of FCU form separate muscle bellies and tendons, and Type III—accessory FCU, where there is an abnormal muscle adjacent to a normal FCU with possible combined features of both FCU and PL. In the current case, there is evidence of an extra muscle in addition to the normal flexor carpi ulnaris, and therefore this is felt to represent a Type III accessory FCU. There are approximately 14 reported cases of Type III FCU anomaly in the anatomy and surgical literature. The embryological basis of the anomaly is thought to be due to the anomalous separation of the flexor blastemal based on the morphology of the reported cases. Based on the previous case reports and proposed embryological hypothesis, there are other potential anomalies associated with the Type III accessory FCU that include an absent PL, duplication of PL and flexor carpi radialis brevis, in addition to variations of the PL such as insertion onto the flexor retinaculum or median nerve innervation.

With regards to the imaging literature, there are several MRI illustrations of the accessory FCU in papers by Lemon et al and Milena et al. Milena et al demonstrated a similar accessory muscle on ultrasound and MRI which also had an insertion to pisiform, as in our case. However, the accessory muscle in their case originated more proximally at the lateral edge of the elbow, blending with the ulnar attachment of pronator quadratus. When considering the accessory muscle origin of our case at the volar/ulnar surface of the distal ulna, Mori et al described four supernumerary muscles of this nature with similar anatomical origin and insertion into the pisiform.

Given the position and location of the accessory muscle, there is potential for ulnar nerve entrapment syndrome with several reports of accessory FCU causing ulnar nerve symptoms that responded well to neurolysis and re-attachment of the distal tendon. In the current case, there were no clinical symptoms of ulnar nerve entrapment. When considering ulnar nerve entrapment syndromes of the upper limb, it should be noted that the accessory anconeus epitrochlearis muscle can cause more proximal ulnar nerve compression in the cubital tunnel, having its origin at the medial epicondyle with insertion to the olecranon. A recent paper found that this accessory muscle accounted for 12 of 142 operated cases of cubital tunnel syndrome.

The ulnar nerve can also be compressed distally at Guyon’s canal. This canal is triangular in shape, with the pisiform forming the medial side of the canal and the deep and superficial volar ligaments forming the deep and superficial boundaries respectively. Within the canal, the ulnar nerve lies medially and the ulnar artery and vein laterally. Reports describe space occupying lesions such as ganglion cyst, lipoma, FCU calcific...
tendonitis and accessory abductor digiti minimi causing ulna nerve symptoms, and our case suggests that hypothetically an accessory FCU insertion towards the distal radial aspect of pisiform could potentially cause reduction in canal volume and subsequent nerve compression.

In conclusion, we have described the ultrasound and MRI appearances of the accessory FCU, a rare muscle variant in the distal forearm which should be considered in the differential diagnosis of distal forearm swellings. Detection of FCU variants is potentially important in the assessment of distal ulnar nerve entrapment syndromes.

**LEARNING POINTS**
- Rare muscle variants in the distal forearm should be considered in the differential diagnosis of distal forearm swellings.
- Detection of FCU variants is potentially important in the assessment of distal ulnar nerve entrapment syndromes.

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Table 1. Summary of Type III accessory FCU (modified from Bhardwaj et al)

| Feature                  | Type III accessory FCU                                                                 |
|--------------------------|--------------------------------------------------------------------------------------|
| In addition to normal FCU; An anomalous muscle in forearm with features of FCU and sometimes PL |
| Due to anomalous separation of flexor blastemal | FCU, flexor carpi ulnaris; PL, palmaris longus. |
| 14                       | Common; absent PL, duplication PL, FCRB                                               |
| Anomalous innervations common | FCRB, flexor carpi radialis brevis; FCU, flexor carpi ulnaris; PL, palmaris longus. |
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