Anatomical Description of a Variant Abductor Digiti Minimi Accessorius Muscle and Its Clinical Correlation with Ulnar Neurovascular Entrapment Syndrome

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Abstract

Variations of the abductor digiti minimi muscle are a rare occurrence as compared to anomalous thenar musculature. Surgico-anatomical basis of such accessory muscular slips should be discussed in detail. This case report presents a rare bilateral existence of accessory slips of abductor digiti minimi originating from the antebrachial fascia and flexor retinaculum in a male cadaver. Hypertrophy of these muscular heads can be occult etiologies in causation of ulnar entrapment neuropathies. Such anomalous muscles can prove to be a boon or bane for an operating surgeon. On one hand, they can be the guiding factor in planning myofascial flaps and on the other; they can cause unwanted iatrogenic complications during antero-medial approach through the wrist tendons. Nevertheless, the presence of accessory bellies in the hypothenar eminence deserve a special mention especially if present bilaterally. The authors have attempted to assign an embryological explanation for the above anomaly and have discussed its possible link with various clinical syndromes.

Keywords: Entrapment, abductor digiti minimi, ulnar nerve, Guyon’s canal, accessory

Introduction

Anomalies of hypothenar musculature are relatively infrequent but rare cases have been reported in the past (1,2). Supernumerary slips of abductor digiti minimi (ADM) have been defined in the past with various terminologies such as abductor longus digiti quinti (3), musculus flexor digiti minimi accessorius (4) and accessorius ad abductorem minimi digitii manus (5). They can arise from the tendons of palmaris longus, flexor carpi radialis, flexor carpi ulnaris, intermuscular septa, flexor retinaculum and rarely from the antebrachial fascia (1,2,3,4,5). This research work reports a rare bilateral existence of accessory ADM arising from the antebrachial fascia with few fibers originating from the flexor retinaculum. Ulnar neuropathy is a rare entity with a wide range of etiologies like ganglion, neoplasm, vascular anomalies, tendinous entrapments and myxoma. Aberrant slips of abductor digiti minimi (ADM) can seldom be the root cause of ulnar entrapment neuropathies (6,7). In clinical practice, these accessory slips can sometimes be asymptomatic and can be encountered during surgical procedures or can sometimes mimic soft tissue tumors (8).

Case Report

Routine anatomic dissection of the manus of a 68-year-old male cadaver revealed the presence of a rare anomalous head of origin for the abductor digiti minimi (ADM) bilaterally. The usual head of ADM originated from the pisiform bone and its distal attachment was on the medial side of the base of the proximal phalanx of the fifth digit, bilaterally.
Figure 1: Figure showing right ADM accessorius. (FCR- flexor carpi radialis; PL- Palmaris longus; FR- flexor retinaculum; AH- accessory head of abductor digiti minimi; FCU- flexor carpi ulnaris; DF- FA- deep fascia of forearm; UA- ulnar artery; UN- ulnar nerve; ADM- abductor digiti minimi; FDM- flexor digiti minimi; PB- Palmaris brevis)

On the right side, an accessory head of the ADM (ADM accessorious) was arising from the ante-brachial fascia of the lower third of the forearm (Fig. 1). Few of the fibers of this accessory muscle also took origin from the flexor retinaculum. This triangular ADM accessorious passed obliquely to the ulnar side of the wrist and traversed through the Guyon’s canal to form a common fused pennate with the normal head of ADM. Few of its fibers also inserted on the pisiform bone. ADM accessorious originated 3.2 cm superior to the proximal border of the flexor retinaculum and was 4.6 cm in length and 2.4 cm broad at its origin. The ulnar neurovascular bundle coursed deep to this accessory head into the Guyon’s canal, the nerve being medial to the artery (Fig. 2). ADM accessorious was related medially to the flexor carpi ulnaris tendon and laterally to the flexor policis longus tendon. This accessory head was lying superficial to the flexor digitorum superficialis. It was innervated by the main trunk of the ulnar nerve. Its arterial supply was derived from the ulnar artery just proximal to the flexor retinaculum.

On the left side, a similar anomalous ADM accessorious originating from the ante-brachial fascia and flexor retinaculum (Fig. 3). It had similar relations as its counterpart and measured 4.2 cm in length and 2.2 cm in breadth. The ulnar neurovascular bundle traversed a course deep to this accessory muscle and supplying it en-route. Morphology and anatomical disposition of other hypothenar muscles was normal, bilaterally.

Discussion

Variations of the ADM are rare but nevertheless have been reported in the past. The commonly reported ones include anomalies like multiple heads of origin, fusion with the flexor digiti minimi brevis, deep abductor-flexor and even its absence (9). Harvie et al. described the incidence of accessory ADM to be as high as 41% with a greater prevalence in males (10). Murata et al. reported the incidence of accessory
ADM with single slips (17%), double slips (80%) and triple slips (3%) (11).

Madhavi and Holla stated that the coexistence of multiple accessory hypothenar muscles is due to their common muscular phylogeny from a single muscle mass (12). The trilaminar theory of hypothenar muscles describes the ADM to be a marginal part of dorsal musculature of hand which has later extended its origin to the ventral aspect (13). It was stated that ADM and flexor digiti minimi are a part of the single muscle representing the last dorsal interosseous (13). ADM has been considered as inner portion of Palmaris longus (13). McMurrich described the developmental existence of flexor digitorum superficialis and suggested that all the hypothenar muscles were its part (14). ADM is one of the earliest muscles to develop in the region of the hypothenar eminence. The growth of its fibers is influenced by the developing ulnar nerve and its arborization. Accessory slips of ADM can be the consequence of variant ulnar developmental patterns (13).

In the present case, the close relation of ADM accessorius to the ulnar neurovascular structures could possibly be responsible for compression neuropathies. The position and bulk of the accessory muscle belly will be deciding factors in aggravating the symptoms such as par aesthesia, paresis, clawing and atrophy (15). Furthermore, muscular spasms release anoxic substances due to anaerobic metabolism and can cause neural irritation and degeneration (16). A rare consequence of ulnar artery entrapment caused by ADM accessorius can be claudicating pain which can mimic peripheral vascular disorders (17). Therefore, vasculopathies involving ulnar artery may result from the ADM accessorious because of its close relation to the ulnar vessels. Hypothenar hammer syndrome is known to be caused by vascular trauma in the hypothenar region, but may also be related to hypertrophied ADM accessorious (18). Intramuscular myxomas usually occur as isolated lesions and are associated with Mazabraud’s or McCune–Albright syndrome (19, 20). Myxomas have been described to occur in the hypothenar muscles and hence ADM accessorious should be borne in mind as an occult site for such myxomas (21).

Abductor digiti minimi rotational flaps have been recently used for soft tissue coverage of the ulnar aspect of the hand and wrist (22). ADM accessorious originating from the fascia and retinaculum as seen in present case may prove to be an excellent alternative for reconstructive surgeries as a vascular pedicle can easily be lifted using the branch from the underlying ulnar artery.

**Conclusion**

Muscular variations as described in the present report having close relation to important neurovascular structures can become symptomatic due to their hypertrophy. Such accessory slips can also be easily harvested for flap reconstructions. Hence, detailed knowledge of such rare variations become imperative for a better clinical approach.

**References**

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