Hypothenar Hammer Syndrome in an Elderly Caucasian Female: A Case Report

Jordan T. Carter 1 , Michael Polmear 1 , Fernando Herrera 2 , Gilberto Gonzalez 1

1. Orthopaedics, Texas Tech University Health Sciences Center, El Paso, USA 2. Plastic Surgery, Medical University of South Carolina, Charleston, USA

Corresponding author: Jordan T. Carter, jordan.carter@ttuhsc.edu

Abstract
Hypothenar hammer syndrome (HHS) is a vascular disorder characterized by ulnar artery thrombosis or aneurysmal formation. It is most commonly reported in middle-aged males who suffer from repetitive trauma to the palm secondary to occupational or recreational activities. Most cases of HHS can be managed conservatively; however, surgical management is typically indicated for persistent symptoms following conservative measures, imminent vascular compromise, those who fail conservative treatment, or those in imminent danger of rupture. Here we present the case of a right-handed 74-year-old female with HHS who presented with a three-week history of a mass in the hypothenar eminence of the left hand. Reportedly, the mass had appeared slowly and had no associated pain, ischemia of the distal hand, numbness, paresthesia, or changes in the overlying skin. She denied any history of a blunt or penetrating hand injury. The patient was treated surgically by removing a 2.0-cm thrombosed pseudoaneurysm of a collateral branch of the ulnar artery within the left hypothenar eminence. On follow-ups at 1, 2, and 12 weeks postoperatively, the patient’s pain was found to be well-controlled. Her normal range of motion was restored, and her digits remained neurovascursively intact. This was an atypical presentation of HHS, and our review of the disorder emphasizes the importance of diagnostic reasoning in rare conditions with unusual presentations of HHS.

Introduction
Hypothenar hammer syndrome (HHS) is a vascular disorder that predominantly affects the dominant hand of middle-aged men who utilize their hand as a “hammer”, arising from repetitive blunt trauma to the hypothenar eminence or, rarely, from a single traumatic blunt injury to the hypothenar eminence [1-5]. This syndrome is caused by the compression and subsequent thrombosis of the ulnar artery against the hamate bone as it passes through Guyon’s canal [1-5]. Patients typically present with complaints of pain, pallor, cyanosis, cold sensitivity, claudication, and color changes localized to the ulnar digits of the affected hand [1-4]. Angiography is usually used to identify the occlusion and define the anatomy, especially if surgical intervention is required. Due to the rarity of this syndrome, no prospective studies have been published so far to identify the most appropriate treatment strategy [1]. Management includes smoking cessation, activity modification, pharmacological therapy, direct thrombin injection, transarterial coiling, and operative intervention for failed conservative management, impending aneurysmal rupture, or ischemia [4-8]. Here we review the literature on HHS and discuss an atypical presentation encountered at our institution.

Case Presentation
We present the case of a right-handed, 74-year-old female who presented with a three-week history of a mass in the hypothenar eminence of the left hand. According to the patient, the mass had appeared slowly and had no associated pain, ischemia of the distal hand, numbness, paresthesia, or changes in the overlying skin. She denied any history of a blunt or penetrating hand injury. The patient was a lifelong nonsmoker and had a past medical history significant for right breast cancer (status: post lumpectomy and radiation), with postoperative right upper extremity lymphedema.

A physical exam of the left upper extremity showed a non-tender, non-pulsatile, firm 1.5 x 1.5-cm subcutaneous mass within the hypothenar eminence at the hook of the hamate (Figure 1). The patient exhibited no neurovascular changes in the hand and demonstrated a normal Allen’s test. Radiographic evaluation showed osteoarthritis of the distal and proximal interphalangeal joints and no acute osseous abnormalities, dislocations, heterotopic ossification, or bone coalitions. An MRI showed a 1.2 x 1.0 x 1.7-cm thrombosed pseudoaneurysm of the distal ulnar artery at the level of Guyon’s canal. Subsequent arterial duplex sonography revealed patent left upper extremity arteries with standard anatomy, including a patent palmar arch.

Keywords: case report, hypothenar hammer syndrome, pseudoaneurysm, surgery, mass

Categories: Cardiac/Thoracic/Vascular Surgery, Plastic Surgery, Orthopedics

Received 12/16/2019
Review began 01/09/2020
Review ended 01/11/2020
Published 01/12/2020

© Copyright 2020
Carter et al. This is an open access article distributed under the terms of the Creative Commons Attribution License CC-BY 3.0., which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

How to cite this article
Carter J T, Polmear M, Herrera F, et al. (January 12, 2020) Hypothenar Hammer Syndrome in an Elderly Caucasian Female: A Case Report. Cureus 12(1): e6636. DOI 10.7759/cureus.6636
The patient was offered pseudoaneurysm resection to prevent possible rupture and ischemia, and she chose to proceed with operative management. During the procedure, the patient was prepped and draped; a tourniquet was inflated to 250 mmHg; an incision was made at Guyon’s canal. A 2.0-cm mass was located radial to the neurovascular bundle, extending from a collateral branch of the ulnar artery after the bifurcation into the deep and superficial branches (Figure 2). The mass was dissected and clipped at its origin, after which the tourniquet was released to confirm that flow through the ulnar artery was intact. The mass was then excised, and the subsequent pathologic examination was consistent with a thrombosed pseudoaneurysm (Figure 3).
FIGURE 2: Intra-operative view
FIGURE 3: Resected pseudoaneurysm

On follow-ups at 1, 2, and 12 weeks postoperatively, the patient presented with the scar seen in Figure 4. Her pain was found to be well-controlled. Her normal range of motion was restored, and her digits remained neurovascually intact.

FIGURE 4: Post-surgical incision

Discussion

Causes

HHS usually results from compression of the ulnar artery just distal to Guyon’s canal [1,3,9]. This course of the ulnar artery narrows at the proximal aspect of the hypothenar eminence where it is surrounded by the
Conclusions

Most cases of HHS are caused by repetitive microtraumas that are thought to stimulate vasospasms causing intimal hyperplasia [12]. This can lead to platelet aggregation, thrombus formation, and occasionally, aneurysmal formation [12,13]. This cycle is propagated as thrombus formation, causing vasospasm of nearby vasculature, which further decreases perfusion [9,12]. This repetitive trauma and insult cause thickening, fibrosis, and scarring [9]. If an aneurysm forms, the nearby ulnar nerve can be compressed, resulting in pain and paraesthesia [1,3,5]. It has been theorized that the limited incidence of HHS, despite a large number of individuals who repetitively use their hand with blunt microtrauma, could be secondary to anatomical variability, subclinical disease, and underlying abnormal arterial histology [1-3,14].

Presentation of hypothenar hammer syndrome

This syndrome is typically seen in laborers, such as carpenters, butchers, and mechanics who use their hand as a "hammer" as part of their profession. HHS has also been associated with sports and hobbies requiring repetitive compression or blunt trauma over the hypothenar eminence. Although rare, HHS has also been documented as resulting from a singular blunt or penetrating trauma to the hypothenar eminence [2,4-7,9,14]. HHS typically presents as pain, pallor, cyanosis, coolness, numbness, and color changes in the ulnar digits secondary to arterial insufficiency [1-5]. Ulceration and gangrene can be present in extreme cases [2]. Occasionally, an associated aneurysm can present as a palpable, pulsatile mass [1,5,4].

HHS is typically observed in the ulnar digits of the dominant hand in males [1,3-7,14]. The thumb is typically spared, allowing for differentiation from Raynaud’s phenomenon [1,4]. The average age of onset is in the fifth decade of life [1,4,7]; however, cases have been reported in older individuals [15]. Most patients cite an average duration of symptoms of two-and-a-half months, highlighting the typical subacute development of HHS [7].

As HHS is likely under-reported, its incidence is hard to determine. One study found 22 cases (1.7%) in 1,300 patients referred to their center for hand ischemia [14]. A larger study found that of the 4,148 patients referred for Raynaud’s phenomenon, 47 (1.1%) cases were actually HHS [1].

Diagnosis

Diagnosis of HHS requires clinical suspicion and can be confirmed via different imaging modalities, such as the characteristic "ying-yang sign" on ultrasound or CT angiography [5,6]. On physical exam, an abnormal Allen’s test [2], suggesting a degree of occlusion in the ulnar artery, and the unilateral nature of HHS helps differentiate it from similarly presenting pathologies, such as Raynaud’s phenomenon [9]. Although HHS can result in aneurysmal formation, the presence of an aneurysm should also prompt consideration of congenital, mycotic, and penetrating trauma etiologies [16].

Catheter-based angiography remains the gold standard to identify the lesion and nearby anatomy, especially if surgical intervention is being considered [1]. CT angiography has high sensitivity (90-95%) and specificity (98-100%) for HHS [1]. Advanced imaging can help to elucidate nearby structures and underlying bony or muscular abnormalities [1,4,7,9,12,17,18]. Regardless of imaging choice, the proximal segments of the nerve must also be examined to ensure that a proximal obstruction is not present [17].

Treatment

Due to the relatively rare nature of HHS, there is no consensus on the best management of the condition [1,3,7]. Conservative means consist of patient education on hand protection, avoiding cold and exacerbating activities and, perhaps most importantly, counseling on smoking cessation. Patients who continue to smoke typically fail both conservative and surgical interventions [5,7]. Pharmacological options include calcium channel blockers, antiplatelet or anticoagulant agents, and pentoxifylline [1,8,10]. Intra-arterial thrombolysis can be offered on the premise of reducing distal embolic events [19]. In one study of four patients with HHS, three experienced improvements after intra-arterial thrombolysis and there were no adverse bleeding events [20]. Conversely, the use of ultrasound-guided thrombin injection has been used with mixed results in the treatment of small vessel pseudoaneurysms including HHS [5,6]. Conservative treatment is usually effective, and a small case series have demonstrated improvement in as many as 83% of patients with conservative treatment [7,8]. However, recurrence is a concern and has been demonstrated in up to 27.7% of conservatively managed cases, namely in patients who continued smoking throughout treatment [6,7]. Surgery is typically indicated for ischemia and vascular damage without adequate collateral circulation [4,6,8]; and options include arterial ligation, resection of the thrombosed or aneurysmal segment with end-to-end anastomosis, or arterial resection with graft reconstruction [1,6-8,10].

Conclusions
HHS is a relatively rare syndrome, typically presenting in middle-aged males who undergo repetitive trauma to the ulnar side of the palm. This case represented a unique presentation of HHS in a 74-year-old female with no history of trauma or repetitive activity to the hypothenar eminence. Also novel to this presentation was the occurrence of a thrombosed pseudoaneurysm in the non-dominant hand and the lack of associated symptoms, leading us to ponder on the etiology of this patient’s condition given the lack of any typical causative factors. This case, in particular, emphasizes the importance of diagnostic reasoning in rare presentations of HSS. Due to the unique presentation seen in the case above, the diagnosis could have been missed at the expense of morbidity to the patient.

**Additional Information**

**Disclosures**

**Human subjects:** Consent was obtained by all participants in this study. **Conflicts of interest:** In compliance with the ICMJE uniform disclosure form, all authors declare the following: **Payment/services info:** All authors have declared that they have no financial support was received from any organization for the submitted work. **Financial relationships:** All authors have declared that they have no financial relationships at present or within the previous three years with any organizations that might have an interest in the submitted work. **Other relationships:** All authors have declared that there are no other relationships or activities that could appear to have influenced the submitted work.

**References**

1. Ablett CT, Hackett LA: Hypothenar hammer syndrome: case reports and brief review. Clin Med Res. 2008, 6:3-8. 10.5212/cmr.2008.775
2. Swanson KE, Bartholomew JR, Paulson R: Hypothenar hammer syndrome: a case and brief review. Vasc Med. 2012, 17:108-115. 10.1177/1358863X11425878
3. Kumar Y, Hossen K, Lu L, Karol I: Ulnar artery aneurysm and hypothenar hammer syndrome. BMJ Case Rep. 2015, 2015:1-2. Accessed: January 12, 2020: https://www.ncbi.nlm.nih.gov/pubmed/26604228. 10.1136/bcr-2015-211444
4. Taj S, Malamis AP, Lomasney LM, Demos TC, Bednar MS: Radiologic case study. Hypothenar hammer syndrome. Orthopedics. 2010, 33:286. 10.3928/01477447-20100329-32
5. Cook JW, Konieczny M, Simons M: The ulnar artery pseudoaneurysm. BMJ Case Rep. 2015, 2015:1-2. Accessed: January 12, 2020: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4611870/. 10.1136/bcr-2015-212791
6. Komorowska-Timek E, Teruya TH, Abou-Zamzam AM Jr, Papa D, Ballard JL: Treatment of radial and ulnar artery pseudoaneurysms using percutaneous thrombin injection. J Hand Surg Am. 2004, 29:956-942. 10.1016/j.jhtra.2004.05.009
7. Marie I, Herve F, Primard E, Cailleux N, Levesque H: Long-term follow-up of hypothenar hammer syndrome: a series of 47 patients. Medicine (Baltimore). 2007, 86:534-543. 10.1097/MD.0b013e3181595d43
8. Lifchez SD, Higgins JP: Long-term results of surgical treatment for hypothenar hammer syndrome. J Reconstr Surg. 2009, 124:210-216. 10.1097/PRS.0b013e318158f7b0
9. Drapel J, Fedy A, Guerini H, Desmarais E, Godefroy D, Le Viet D, Chevrier A: Vascular lesions of the hand. Eur J Radiol. 2005, 56:351-354. 10.1016/j.ejrad.2005.03.013
10. McCready RA, Bryant MA, Divelbiss JL: Combined thenar and hypothenar hammer syndromes: case report and review of the literature. J Vasc Surg. 2008, 48:741-744. 10.1016/j.vxs.2008.05.055
11. Corn J Jr, Bergan JF, Bell JL: Hypothenar hammer syndrome: posttraumatic digital ischemia. Surgery. 1970, 68:1122-1128.
12. Genchellia H, Demir MK, Unlu E, Temizöz O, Ozdemir H: Hypothenar hammer syndrome: gray-scale and color Doppler sonographic appearances. J Clin Ultrasound. 2008, 36:98-100. 10.1002/jcu.20564
13. Abudakka M, Pillai A, Al-Khaif A: Hypothenar hammer syndrome: rare or underdiagnosed?. Eur J Vasc Endovasc Surg. 2006, 32:257-260. 10.1016/j.ejvs.2006.02.013
14. Ferris BL, Taylor LM Jr, Gomaya K, McLaugherty RB, Edwards JM, Moneta GL, Porter JM: Hypothenar hammer syndrome: proposed etiology. J Vasc Surg. 2000, 31:104-115. 10.1016/s0741-5214(00)00772-3
15. Carter PM, Hollinshead PA, Desmond JS: Hypothenar hammer syndrome: case report and review. Clin Med Res. 2008, 6:3-8. 10.1016/j.cmr.2008.05.055
16. Cooke RA: Hypothenar hammer syndrome: a discrete syndrome to be distinguished from hand-arm vibration syndrome. Occup Med (Lond). 2003, 53:320-324. 10.1093/occmed/koq071
17. Torre J: Ulnar artery aneurysm with digital ischemia. Vasc Med. 1999, 4:143-145. 10.1177/1358863X990040204
18. Abdel-Gawad EA, Bonatti H, Housseini AM, Maged IM, Morgan RF, Hagspiel KD: Hypothenar hammer syndrome in a computer programmer: CTA diagnosis and surgical and endovascular treatment. Vasc Endovascular Surg. 2009, 43:509-512. 10.1177/153874400934346
19. Bakhach J, Chahidi N, Conde A: Hypothenar hammer syndrome: management of distal embolization by intra-arterial fibrinolitics. Chir Main. 1998, 17:215-220. 10.1016/s0755-9653(98)80804-1
20. Wheatley MJ, Marx MV: The use of intra-arterial urokinase in the management of hand ischemia secondary to palmar and digital arterial occlusion. Ann Plast Surg. 1996, 37:556-562. 10.1097/00000637-199610000-00005