Atypical Presentation of Right Ventricular Cardiac Hamartoma in a Young Man

RAMEZ BARSOOM, MS
LAMEES I. EL NIHUM, MD, MENG
QASIM AL ABRI, MD
AREEBA ALI, MD
SUSAN L. HALEY, MD

MOHAMMED A. CHAMSI-PASHA, MD
MORITZ C. WYLER VON BALLMOOS, MD, PHD, MPH
THOMAS E. MACGILLIVRAY, MD
MICHAEL J. REARDON, MD

*Author affiliations can be found in the back matter of this article

ABSTRACT

Cardiac tumors in adults are exceedingly rare and usually benign. We describe a 29-year-old man with a previous diagnosis of interventricular septal hypertrophy who presented with increasing severity of dyspnea and fatigue. Work-up revealed a 4.9 × 3.7 cm mass at the base of the interventricular septum. Biopsy revealed a benign cardiac hamartoma atypically located in the right ventricle, and the mass was resected via right ventriculotomy.

CORRESPONDING AUTHOR:
Michael J. Reardon, MD
Methodist DeBakey Heart & Vascular Center, Houston Methodist Hospital, Houston, Texas, USA
mreardon@houstonmethodist.org

KEYWORDS:
cardiac hamartoma; ventriculotomy; cardiac tumor; echocardiography; mass; right ventricle

TO CITE THIS ARTICLE:
Barsoom R, El Nihum LI, Al Abri Q, Ali A, Haley SL, Chamsi-Pasha MA, von Ballmoos MCW, MacGillivray TE, Reardon MJ. Atypical Presentation of Right Ventricular Cardiac Hamartoma in a Young Man. Methodist DeBakey Cardiovasc J. 2022;18(1):102-107. doi: 10.14797/mdcvj.1158
INTRODUCTION

Hamartoma of mature cardiac myocyte is a relatively recent term classified under benign tumor and tumor-like lesions of the heart. They often present a diagnostic challenge due to their nonspecific symptoms, which can include chest pain, fatigue, dyspnea, or palpitations. The present case has been highlighted due to its rarity, unique infiltrative radiological appearance, and poorly defined clinicopathologic spectrum of features.

CASE REPORT

A 29-year-old male with a previous diagnosis of hypertrophic cardiomyopathy presented to his cardiologist for episodes of fatigue, chest pain, and progressive dyspnea on physical exertion for the past 3 years, increasing in frequency and severity over the last 6 months. During this time, he experienced several episodes of ventricular tachycardia and one episode of near syncope. He was diagnosed with hypertrophic cardiomyopathy 4 years prior to presentation after a murmur was heard on routine exam, and it had been medically managed with a beta blocker.

Transthoracic echocardiogram revealed a prominent thickening at the base of the interventricular septum that extended into the right ventricular outflow tract (RVOT) (Figure 1 A-C). It was noncontractile and heterogeneous in appearance with a vascular component, suggestive of a tumor. The mass was confirmed on subsequent cardiac magnetic resonance imaging, which demonstrated a 4.9 × 3.7 cm mass extending from the anterior left ventricular wall to the inferoseptum and from the base of the heart through the mid-segment as well as into the RVOT (Figure 2 A-D). Cardiac computed tomography showed a basal anteroseptal mass protruding into the RVOT with central contrast enhancement and multiple septal perforator feeding vessels (Figure 2 E). At this time, the differential diagnosis included both benign and malignant etiologies.

A fluorodeoxyglucose positron emission tomography scan was then performed and demonstrated an area of hypermetabolism in the anterior aspect of the right ventricle medially. The maximum standardized uptake value (SUVmax) of this abnormality was difficult to assess due to the proximity of the left myocardium, which is normally and physiologically hypermetabolic. SUVmax of the lesion was approximated to be 7.0, concerning for the presence of malignant tissue. Given the vascular nature of the mass, a coronary angiogram and biopsy were performed in anticipation of surgical planning. Cardiac catheterization revealed normal coronary arteries and disorganized vascularization towards the septum. Right ventricular mass biopsy showed no neoplasia; however, there was some innocuous subendocardial fibrous tissue. Pathology report showed hypertrophic cardiomyocytes consistent with a cardiac hamartoma.

Due to significant symptoms with minimal exertion secondary to the mass causing RVOT obstruction, a palliative resection was performed via a right ventriculotomy. Postoperative pathology of the specimen showed a soft, tan, flesh-colored mass measuring 3 × 1.6 × 1.4 cm. Sections showed discrete lesions of marked myocyte hypertrophy with disorganization, focal scarring, and thickened intramural arteries (Figure 3 A-C).

DISCUSSION

Hamartomas consisting of mature cardiac myocytes are rare; as such, their clinicopathologic spectrum is not well defined in the literature. Based on a literature search, only 30 patients with a diagnosis of hamartoma of mature cardiac myocytes have been identified from 1998 to 2020 (Table 1). Of these 30, 19 were males...
(63%) and 11 were females (37%). The mean age was 32 ± 21 years, ranging from 6 months to 76 years. Eight of the 30 patients (27%) were identified in the pediatric population. The hamartoma was localized to the left ventricle in 13 patients, the right atrium in 7 patients, and the right ventricle in 3 patients; 2 patients showcased multiple locations. Ten of these patients (33%) were asymptomatic at time of presentation. The present case was an atypical presentation of hamartoma due to its location in the right ventricle rather than in the free wall of the left ventricle, which is the most commonly reported.

Figure 2 Preoperative cardiac imaging. Magnetic resonance imaging using steady state free precession cine sequence: (A) 3-chamber view, (B) short axis at the mid-level, (C), first pass perfusion, and (D) delayed enhancement after gadolinium showing a large right ventricular (RV) mass measuring 4.9 × 3.7 cm invading the RV free wall, with (E) intense first pass perfusion and significant late gadolinium enhancement cardiac computed tomography demonstrating short axis basal anteroseptal mass protruding into the RV outflow tract with central contrast enhancement and multiple septal perforator feeding vessels.

Figure 3 H&E stain at increasing magnification 100×. (A) The resected lesion was composed of disorganized, hypertrophic cardiac myocytes with interstitial fibrosis. (B) There were also scattered thickened intramural arteries, dilated venules, and small collections of adipocytes 200×. (C) At intermediate magnification, the myocytes were haphazardly arranged and enlarged with sarcoplasmic vacuolation and nuclear enlargement 400×. There were scattered enlarged nuclei, some with mild hyperchromasia, irregular nuclear contours, and inconspicuous nucleoli.
| YEAR | PUBLISHED | AUTHOR | AGE | GENDER | CLINICAL SYMPTOMS | DIAGNOSTIC EVALUATION | TUMOR LOCATION | TUMOR SIZE |
|------|-----------|--------|-----|--------|-------------------|-----------------------|----------------|-----------|
| 1998 |           | Sturtz et al. | 24  | Male   | Hypertension, palpitations, and premature contractions | —                     | Left ventricle   | —         |
| 1998 |           | Burke et al.  | 22  | Male   | Asymptomatic      | Echocardiogram         | Right ventricle  | 5 cm      |
| 1998 |           | Burke et al.  | 28  | Male   | WPW syndrome and an episode of syncope | Echocardiogram         | Right atrium    | —         |
| 1998 |           | Burke et al.  | 9   | Male   | Sudden death      | Autopsy                | Right atrium    | 1 × 2 cm  |
| 2001 |           | Dinh et al.   | 33  | Male   | Generalized tachycardia | Echocardiogram         | Left ventricle  | 4.5 × 3.1 × 4.4 cm |
| 2004 |           | Chu et al.    | 76  | Male   | History of hypertension | In surgery             | Crista terminalis | 0.5 × 1 × 0.5 cm |
| 2005 |           | Martinez QM et al. | 33  | Male   | Palpitations and dyspnea | Echocardiogram         | Left ventricle  | 4.5 × 5.5 cm |
| 2008 |           | Movahedi et al. | 58  | Male   | Progressive dyspnea | In surgery             | Right atrium    | 1.5 × 1 × 0.5 cm |
| 2008 |           | Fealey et al. | 0.5 | Male   | Asymptomatic      | —                     | —              | —         |
| 2008 |           | Fealey et al. | 0.5 | Female | Asymptomatic      | —                     | —              | —         |
| 2008 |           | Fealey et al. | 1.2 | Male   | Asymptomatic      | —                     | —              | —         |
| 2008 |           | Fealey et al. | 10  | Male   | Asymptomatic      | Echocardiogram         | Left ventricle  | 5 × 3 cm  |
| 2008 |           | Fealey et al. | 16  | Female | Asymptomatic      | Echocardiogram         | Right ventricle  | 8 × 9 cm  |
| 2008 |           | Fealey et al. | 57  | Male   | Sudden death      | —                     | —              | —         |
| 2008 |           | Fealey et al. | 74  | Male   | Exertional dyspnea | —                     | —              | —         |
| 2008 |           | Menon et al.  | 16  | Female | Weight loss       | Echocardiogram         | Right ventricle  | 8 × 7 × 3 cm |
| 2008 |           | Menon et al.  | 10  | Male   | Asymptomatic      | Echocardiogram         | Left ventricle  | 0.3 × 0.2 cm |
| 2009 |           | Hsu et al.    | 19  | Female | Intermittent palpitations and dizziness | Echocardiogram         | Left ventricle  | 4 × 7 cm  |
| 2009 |           | Galeone et al. | 56  | Female | Asymptomatic      | Chest CT               | Pulmonary infundibulum | 9 × 9 × 4 cm |
| 2011 |           | Dell’Amore et al. | 35  | Female | Palpitations and dyspnea | Echocardiogram         | Left ventricle  | 4.2 × 3.3 × 2.7 cm |
| 2013 |           | Raffa et al.  | 41  | Female | Chest pain        | Echocardiogram         | Right atrium    | 2.5 × 1.3 cm |
| 2017 |           | Ayaoub et al. | 14  | Male   | Asymptomatic      | Echocardiogram         | Left ventricle  | 9 × 5 × 6 cm |
| 2017 |           | Hadravská et al. | 39  | Female | Ruptured aneurysm and severe pneumonia | Autopsy | Left ventricle | 4.5 × 3 × 3 cm |
| 2017 |           | Abuzaid et al. | 21  | Female | Chest pain and dyspnea | Echocardiogram         | Left ventricle  | 1.6 × 1.3 × 1.9 cm |
| 2017 |           | Liu et al.    | 64  | Male   | Dyspnea           | Echocardiogram         | Left ventricle  | 2 × 3 cm  |
| 2018 |           | Mantilla-Hernandez et al. | 23  | Female | Paroxysmal nocturnal dyspnea, RVR and edema | Autopsy | Right atrium | —         |
| 2018 |           | Negri et al.  | 44  | Female | Asymptomatic      | Echocardiogram         | Left ventricle  | 4 × 3 cm  |
CONCLUSION

Hamartomas prove to be a diagnostic challenge due to nonspecific and often nonexistent symptoms. Given this as well as their slow-growing nature, surgical resection followed by postoperative microscopic pathological examination is the only reliable method of definitive diagnosis. However, microscopic features including myocyte hypertrophy and interstitial fibrosis in a whorled pattern are also nonspecific and characteristic of hypertrophic cardiomyopathy as well, which was the initial diagnosis of our patient. Distinction between hamartomas and hypertrophic cardiomyopathy on a small surgical or biopsy specimen has been proven difficult and inconclusive at times. Additionally, imaging studies alone are not pathognomonic for hamartoma. Thus, establishing a diagnosis requires a combination of clinical findings, imaging studies, and microscopic examinations.

COMPETING INTERESTS

Michael J. Reardon, MD, is a consultant for Medtronic, Boston Scientific, and Gore Medical. Moritz C. Wyler von Ballmoos, MD, PhD, MPH, has a financial relationship with Medtronic and Boston Scientific. All other authors have no competing interests to declare.

AUTHOR AFFILIATIONS

Ramez Barsoom, MS orcid.org/0000-0002-6287-4753
Houston Methodist DeBakey Heart & Vascular Center, Houston Methodist Hospital, Houston, Texas, USA; Texas A&M University School of Medicine, Bryan, Texas, USA

Lamees I. El Nihum, MD, MEng
Houston Methodist DeBakey Heart & Vascular Center, Houston Methodist Hospital, Houston, Texas, USA; Texas A&M College of Medicine, Bryan, Texas, USA

Qasim Al Abri, MD
Houston Methodist DeBakey Heart & Vascular Center, Houston Methodist Hospital, Houston, Texas, USA

Areeba Ali, MD orcid.org/0000-0002-5694-1753
Houston Methodist DeBakey Heart & Vascular Center, Houston Methodist Hospital, Houston, Texas, USA

Susan L. Haley, MD orcid.org/0000-0001-5942-1705
Department of Pathology & Genomic Medicine, Houston Methodist Hospital, Houston, Texas, USA

Mohammed A. Chamsi-Pasha, MD orcid.org/0000-0002-1445-4789
Houston Methodist DeBakey Heart & Vascular Center, Houston Methodist Hospital, Houston, Texas, USA

Moritz C. Wyler von Ballmoos, MD, PhD, MPH
Houston Methodist DeBakey Heart & Vascular Center, Houston Methodist Hospital, Houston, Texas, USA

Thomas E. MacGillivray, MD
Houston Methodist DeBakey Heart & Vascular Center, Houston Methodist Hospital, Houston, Texas, USA

Michael J. Reardon, MD orcid.org/0000-0002-2880-6132
Houston Methodist DeBakey Heart & Vascular Center, Houston Methodist Hospital, Houston, Texas, USA

REFERENCES

1. Sturtz CL, Abt AB, Leuenberger UA, Damiano R. Hamartoma of mature cardiac myocytes: a case report. Mod Pathol. 1998 May;11(5):496-9. PMID: 9619605

2. Burke AP, Ribe JK, Bajaj AK, Edwards WD, Farb A, Virmani R. Hamartoma of mature cardiac myocytes. Hum Pathol. 1998 Sep;29:904-9. doi: 10.1016/s0046-8177(98)90194-0

3. Dinh MH, Galvin JM, Aretz TH, Torchiana DF. Left ventricular hamartoma associated with ventricular tachycardia. Ann Thorac Surg. 2001 May;71(5):1673-5. doi: 10.1016/s0003-4975(00)02312-2

4. Chu PH, Yeh HI, Jung SM, et al. Irregular connexin43 expressed in a rare cardiac hamartoma containing adipose tissue in the crista terminalis. Virchows Arch. 2004 Apr;444(4):383-6. doi: 10.1007/s00428-003-0960-6

5. Martínez Quesada M, Trujillo Berraquero F, Almendro Delia M, Hidalgo Urbano R, Cruz Fernández JM. [Cardiac hamartoma. Case report and literature review]. Rev Esp Cardiol. 2005 Apr;58(4):450-2. PMID: 15847741
6. Movahedi N, Boroumand MA, Sotoudeh AM, Yazdanifard P. Mature cardiac myocyte hamartoma in the right atrium. Asian Cardiovasc Thorac Ann. 2008 Oct;16(5):e47-8. doi: 10.1177/021849230801600525

7. Fealey ME, Edwards WD, Miller DV, Menon SC, Dearani JA. Hamartomas of mature cardiac myocytes: report of 7 new cases and review of literature. Hum Pathol. 2008 Jul;39(7):1064-71. doi: 10.1016/j.humpath.2007.11.014

8. Menon SC, Miller DV, Cabalka AK, Hagler DJ. Hamartomas of mature cardiac myocytes. Eur J Echocardiogr. 2008 Nov;9(6):835-9. doi: 10.1093/ejehocard/jen182

9. Hsu PS, Chen JL, Hong GJ, Tsai YT, Tsai CS. Heart transplantation for ventricular arrhythmia caused by a rare hamartoma. J Heart Lung Transplant. 2009 Oct;28(10):1114-5. doi: 10.1016/j.healun.2009.06.015

10. Galeone A, Valdile P, Gayet JB, Laborde F. Hamartoma of mature cardiac myocytes of the pulmonary infundibulum. Interact Cardiovasc Thorac Surg. 2009 Dec;9(6):1029-31. doi: 10.1510/icvts.2009.215855

11. Dell’Amore A, Lanzanova G, Silenzio A, Lamarras M. Hamartoma of mature cardiac myocytes: case report and review of the literature. Heart Lung Circ. 2011 May;20(5):336-40. doi: 10.1016/j.hlc.2011.01.015

12. Raffa GM, Malvindi PG, Settepani F, et al. Hamartoma of mature cardiac myocytes in adults and young: case report and literature review. Int J Cardiol. 2013 Feb 20;163(2):e28-30. doi: 10.1016/j.ijcard.2012.08.052

13. Ayoub C, Luis SA, Maleszewski JJ, Pellikka PA. Advanced cardiac imaging techniques assist in characterizing a cardiac mass and directing management. Echocardiography. 2017 Nov;34(11):1744-1746. doi: 10.1111/echo.13719

14. Hadravská Š, Dubová M, Miesbauerová M, et al. [Hamartoma of mature cardiac myocytes. Autopsy case report]. Cesk Patol. 2017 Jan;1;53(4):185-187. PMID: 29227122

15. Abuzaid AS, Gakhal M, Montgomery E, LaPoint R, Horn R, Banbury MK. Cardiac Hamartoma: A Diagnostic Challenge. CASE (Phila). 2017 Apr 24;1(2):59-61. doi: 10.1016/j.case.2017.01.014

16. Liu L, Qin C, Guo Y. Rare case of left ventricular mesenchymal hamartoma. J Thorac Cardiovasc Surg. 2018 Jan;155(1):346-350. doi: 10.1016/j.jtcvs.2017.09.051

17. Mantilla-Hernández JC, Amaya-Mujica J, Alvarez-Ojeda OM. An unusual tumour: Hamartoma of mature cardiac myocytes. Rev Esp Patol. 2019 Jan-Mar;52(1):50-53. doi: 10.1016/j.patol.2018.07.004

18. Negri F, De Luca A, Aniello P, et al. Cardiac hamartoma. J Card Surg. 2018 Oct;33(10):640-642. doi: 10.1111/jocs.13802

19. Kumari K, Arava S, Kumar S, Bansal A, Bisoi AK, Ray R. Hamartoma of mature cardiac myocytes: Report of a rare case with review of literature. J Pract Cardiovasc Sci. 2018;4(1):55-58. doi: 10.4103/jpcs.jpcs_15_18

20. Fu F, She X. Mature cardiac myocytohamartoma: a case report and review of literature. Int J Clin Exp Pathol. 2019;12(4):1424-1428. Published 2019 Apr 1. PMID: 31933959

21. Zhou X, Zhou Y, Zhaoshun Y, et al. Hamartoma of mature cardiomyocytes in right atrium: A case report and literature review. Medicine (Baltimore). 2019 Aug;98(31):e16640. doi: 10.1097/MD.0000000000016640

TO CITE THIS ARTICLE:
Barsoom R, El Nihum LI, Abri A, Ali A, Haley SL, Chamsi-Pasha MA, von Ballmoos MCW, MacGillivray TE, Reardon MJ. Atypical Presentation of Right Ventricular Cardiac Hamartoma in a Young Man. Methodist DeBakey Cardiovasc J. 2022;18(1):102-107. doi: 10.14797/mdcvj.1158

Submitted: 03 August 2022   Accepted: 22 September 2022   Published: 04 October 2022

COPYRIGHT:
© 2022 The Author(s). This is an open-access article distributed under the terms of the Attribution-NonCommercial 4.0 International (CC BY-NC 4.0), which permits unrestricted use, distribution, and reproduction in any noncommercial medium, provided the original author and source are credited. See https://creativecommons.org/licenses/by-nc/4.0/.

Methodist DeBakey Cardiovascular Journal is a peer-reviewed open access journal published by Houston Methodist DeBakey Heart & Vascular Center.