Pseudoaneurysm Arising from Mitral Aortic Intervalvular Fibrosa (P-MAIVF) Communicating with Left Atrium (LA): Multiple Detector Computed Tomography (MDCT) Evaluation

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Summary

Background: The entity pseudoaneurysm arising from the mitral aortic intervalvular fibrosa (P-MAIVF) is a rare cardiac finding caused by multiple factors. This entity is usually diagnosed with echocardiography and confirmed with cardiac computed tomography (CT).

Case Report: We presented a case of congenital P-MAIVF communicating with the left atrium (LA) and an aberrant right subclavian artery, misdiagnosed as primary mitral regurgitation (MR) in transthoracic echocardiogram (TTE) due to relative contraindications to transesophageal echocardiogram (TEE), revealed in a hemophilic patient, and diagnosed with cardiac CT.

Conclusions: In conclusion, cardiac CT plays a definitive role not only in anatomical assessment and confirmation of the lesion but also in primary diagnostics in patients suspected of MAIVF – especially those with relative and absolute contraindications to TEE.

MeSH Keywords: Aneurysm, False • Congenital Abnormalities • Hemophilia A

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Background

P-MAIVF is a unique and rare entity especially for a radiologist as it arises from a complex anatomical area in the aorto-mitral continuity region. P-MAIVF is caused by multiple factors grouped into surgical, infective, inflammatory and congenital ones [1,2]. Variable symptoms and dreaded complications are associated with an enlarging aneurysm [3]. The primary screening investigation is TTE and it is complemented by TEE for diagnostic purposes. Confirmation and anatomical assessment are carried out with the use of MDCT and Cardiac Magnetic Resonance Imaging (MRI), which is useful for surgical evaluation [4]. The recommended treatment for P-MAIVF is surgery in both asymptomatic and symptomatic patients [5].

Case Report

A 29-year-old hemophilic male patient was admitted to the cardiology unit of our hospital with complaints of New York Heart Association (NYHA) Grade II dyspnea for three years. There was no past surgical history. On auscultation there was holosystolic murmur with a diastolic component. A provisional diagnosis of mitral regurgitation was made. He had been investigated with TTE in various hospitals for his symptoms, with a consistent diagnosis of primary MR. There was no past history of any ischemic heart disease, rheumatic or infective heart condition. The patient was referred to us for cardiac CT in view of persistent symptoms.

Cardiac CT was performed with a Brilliance™ 64-slice scanner (Philips Medical Systems, Cleveland, Ohio, USA) using an adult cardiac CT protocol and retrospective electrocardiogram gating. A volume of 100 mL of iodinated contrast medium was administered, followed by saline solution. The images were processed on a dedicated Aquarius™ Workstation (TeraRecon, San Mateo, Calif.); reformatted MPR and MIP images in different planes were analyzed.
Cardiac CT revealed an out-pouching coming posteriorly from the mitral aortic intervalvular fibrosa (MAIVF). The apex of the aneurysm was separated from the ascending aorta by the pulmonary trunk. The left coronary artery was abutting the aneurysm anteriorly. The left atrium and the left ventricle were both enlarged and the aneurysm had a fistulous communication with the left atrium (LA). Calcification was also noted in the wall of the aneurysm. There was an associated aberrant right subclavian artery (AbnSCA) from the aorta (Ao), having retrotracheal course. SVC indicates the superior vena cava, TR indicates the trachea.

The absence of spikes of fever, negative blood cultures, normal C-reactive protein level and negative Anti-streptolysin O antibody test result ruled out infective endocarditis as an aetiologic factor. Normal erythrocyte sedimentation rate and aortogram also ruled out the inflammatory cause such as Takayasu’s arteritis, which is a possible etiological factor of pseudoaneurysms.

The patient was subsequently operated on and the findings of cardiac CT were confirmed. The histopathological report
revealed the fibrocollagenous nature of the wall along with calcifications suggestive of a pseudoaneurysm. No signs of inflammatory activity were noted. Culture of the excised tissue did not yield any growth. Thus, a final diagnosis of congenital pseudoaneurysm of MAIVF communicating with LA along with an aberrant right subclavian artery was made.

Discussion

Congenital subaortic aneurysms are rare. They were initially described in the African population and subsequently reported on in other racial groups. Many of the subaortic aneurysms arise from MAIVF. A comprehensive review study of articles published between 1966–2009 revealed 90 patients with a pseudoaneurysm of MAIVF out of which 9 cases had communication of the pseudoaneurysm with LA [1].

Left and noncoronary leaflets of the aortic valve share fibrous continuity with the anterior/aortic leaflet of the mitral valve. The thick ends of this fibrous area continue with the ventricular musculature on both sides and are called right and left fibrous trigones. The MAIVF is a central triangular area bounded by these right and left trigones [6].

The causes of P-MAIVF are grouped into surgical, infective, inflammatory and congenital. The most common etiology is the aortic valve surgery and infective endocarditis [1]. Subaortic aneurysms have also been associated with tuberculosis, syphilis, rheumatic carditis and Takayasu’s arteritis [7]. Some of these lesions are thought to be congenital in origin [2]. Being a relatively avascular area, it is prone to weakening and abscess collection after aortic [8,9] and, less commonly, mitral valve surgeries [10,11].

Infective endocarditis of the aortic valve leads to infection of MAIVF. Infection of this area leads to the formation of a subaortic abscess or pseudoaneurysm of the left ventricular outflow tract (LVOT). Therefore, a pseudoaneurysm can rupture in the left atrium due to systolic jet from the left ventricular outflow tract [12].

A patient can be asymptomatic or present with dyspnea, infective symptoms (due to endocarditis), angina (due to compression of coronary arteries) and congestive cardiac failure. Various complications of pseudoaneurysms of MAIVF have been reported [1]. Enlargement of a pseudoaneurysm may lead to compression of adjacent structures including the left atrium, coronary arteries [3], and the pulmonary artery. Proximity to the left atrium and aorta may result in fistulous communications with these structures (Figure 3). Other reported complications include rupture into the pericardial space [13]. Mitral annulus dysfunction may also lead to functional mitral regurgitation [14].

The first screening investigation to start with is TTE. However, due to posterior location of pseudoaneurysms, TEE is preferred, and communication with LA can mislead the diagnosis to primary MR in TTE, as in our case [1]. A golden standard to confirm the diagnosis is catheter angiography. With an advent of MDCT and cardiac MRI, invasive catheterization is unnecessary. CT angiography has the advantage of simultaneous evaluation of valvular structures, coronary arteries, thrombi, and of a precise location of a pseudoaneurysm with regard to other cardiac chambers, aortic root and pulmonary artery. Transesophageal echo has been used extensively to demonstrate pseudoaneurysms of MAIVF. However, to the best of our knowledge, the use of cardiac CT to delineate the anatomy of P-MAIVF is rare; only a few cases have been reported on with the use of CT [4,9,11,15–19].

The recommended treatment for P-MAIVF is surgery in both asymptomatic and symptomatic patients. The patch repair in P-MAIVF can be carried out with or without aortic root reconstruction. Percutaneous closure of a pseudoaneurysm can also be performed in a candidate surgically not suitable [5].

We presented this unusual and rare case of P-MAIVF communicating with LA (which is never reported on in literature on cardiac CT imaging), with an aberrant right subclavian artery, undiagnosed for several years. Our patient, due to coagulopathy, never underwent TEE, and was diagnosed as primary MR on TTE for years. Pseudoaneurysm of MAIVF is a rare cardiac finding caused by multiple factors and is diagnosed with echocardiography and confirmed with cardiac CT.

Conclusions

We presented a case of a hemophilic patient with a congenital pseudoaneurysm of MAIVF communicating with LA and an aberrant right subclavian artery, diagnosed with the use of cardiac CT, initially misdiagnosed as primary MR on TTE due to relative contraindications to TEE. Cardiac CT plays a definitive role not only in anatomical assessment and confirmation of the lesion type but also in primary diagnostics in patients suspected of MAIVF, especially those with relative and absolute contraindications to TEE.

References:

1. Sudhakar S, Sewani A, Agrawal M et al: Pseudoaneurysm of the mitral-aortic intervalvular fibrosa (MAIVF): A comprehensive review. J Am Soc Echocardiogr; 2010; 23(10): 1009–18
2. Sivasankaran S, Kannan BR, Kumar A et al: Coexistence of congenital subaortic and sinus of valsalva aneurysms. Indian Heart J; 2002; 54: 432–34
3. Bier AJ, Lamphere JA, Daily PO: Coronary artery compression caused by a large pseudoaneurysm of the mitral-aortic intervalvular fibrosa. J Am Soc Echocardiogr; 1995; 8(5 Pt 1): 753–56
4. Ghersin E, Litmanovich D, Agmon Y et al: Pseudoaneurysm of the mitral-aortic intervalvular fibrosa following aortic valve replacement – diagnosis and dynamic evaluation with multidetector CT and transesophageal echocardiography. Interact Cardiovasc Thorac Surg; 2005; 46(6): 502–4
5. Jiménez Valero S, García E, González Pinto A et al: Percutaneous closure of pseudoaneurysm of the mitral-aortic intervalvular fibrosa. Rev Esp Cardiol; 2005; 58(12): 1473–75 [in Spanish]
6. Joshi SS, Jagadeesh AM, Furtado A et al: Transesophageal echocardiography in surgical management of pseudoaneurysm of mitral-aortic intervalvular fibrosa with aneurysms of right sinus of Valsalva and left main coronary artery. Ann Card Anaesth, 2013; 16: 40–43

7. Tufekcioğlu O, Özlu ME, Cay S et al: Pseudoaneurysm of the mitral-aortic intervalvular fibrosa in a patient with Takayasu’s arteritis. Can J Cardiol, 2008; 24: 718

8. Aoyagi S, Fukunaga S, Otsuka H et al: Left ventricular outflow tract pseudoaneurysm after aortic valve replacement: case report. J Heart Valve Dis, 2004; 13(1): 145–48

9. Entrikin DW, Shroff GS, Kon ND et al: Pseudoaneurysm of the mitral-aortic intervalvular fibrosa: a delayed complication of aortic root replacement. J Cardiovasc Comput Tomogr, 2011; 5(5): 333–35

10. Namboodiri N, Dorn SK, Thomas B et al: Subannular left ventricular pseudoaneurysm following mitral valve replacement. J Cardiothorac Surg, 2008; 3: 28

11. Choh NA, Shaheen F, Rather H et al: Pseudoaneurysm of mitral-aortic intervalvular fibrosa in a child: Demonstration by MDCT and MRI. Ann Pediatr Cardiol, 2013; 6(1): 80–82

12. Bansal RC, Graham BM, Jutzy KR et al: Left ventricular outflow tract to left atrial communication secondary to rupture of mitral-aortic intervalvular fibrosa in infective endocarditis: diagnosis by transesophageal echocardiography and color flow imaging. J Am Coll Cardiol, 1990; 15(2): 499–504

13. Ozilbash AH, Schwartz CJ: False aneurysm of left ventricle due to perforation of mitral-aortic intervalvular fibrosa with rupture and cardiac tamponade. Rare complication of infective endocarditis. Am J Cardiol, 1973; 32(1): 110–13

14. Espinosa-Caliani JS, Montijano A, Martíàa Melero JA et al: Pseudoaneurysm in the mitral-aortic intervalvular Fibrosa. A cause of mitral regurgitation. Eur J Cardiothorac Surg, 2000; 17(6): 757–59

15. Yokoyama Y, Tamaki S, Kato N et al: Pseudoaneurysm from the mitral-aortic intervalvular fibrosa following endocarditis. Jpn J Thorac Cardiovasc Surg, 2003; 51(8): 374–77

16. Linhartová K, Veselka J, Adla T: Left ventricular pseudoaneurysm as a late complication of mitral annuloplasty. Eur Heart J, 2007; 28(19): 2360

17. Berrizbeitia LD, Anderson WA: Ultrafast computed tomography in infectious pseudoaneurysm of the left ventricular outflow tract. J Thorac Cardiovasc Surg, 1997; 114(1): 138–39

18. Acioli Pereira L, Pontes Gonçalo E, Alcântara Farran J et al: Giant pseudoaneurysm of the left ventricular outflow tract: a rare disease. Rev Port Cardiol, 2013; 32(6): 541–44

19. Thai IC, Fu YC, Lin PC et al: MDCT evaluation of congenital mitral-aortic intervalvular fibrosa aneurysm: implications for the etiology and differential diagnosis. Pediatr Radiol, 2009; 39(1): 80–83