Cross-sectional imaging of sinus of Valsalva aneurysms: lessons learned

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ABSTRACT

Sinus of Valsalva aneurysm, dilatation of one or more of the aortic sinuses, is a rare but important aortic root defect, which can be a cause of some serious cardiac sequels. The purpose of this article is to review the etiopathogenesis, relevant anatomy, clinical manifestations, potential complications, multimodality imaging features, and management of this rare but important entity of sinus of Valsalva.

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inus of Valsalva (SOV) aneurysm is defined as either the dilatation of one or more of the aortic sinuses located between the aortic valve annulus and the sinotubular junction or a crescentic/tubular-shaped outpouching extending from the body or apex of a normally sized SOV (1–3).

Given the various appearances and difficult location, detection of ruptured and unruptured SOV aneurysms can be a diagnostic challenge. Although, the ruptured SOV aneurysm can be potentially fatal, their prognosis after treatment is excellent. This emphasizes the need for timely accurate diagnosis. In this article, we review the relevant etiopathogenesis, anatomy, clinical manifestations, potential complications, cross-sectional imaging features, and management of this rare but important aortic root defect.

SOV aneurysms prevalence and etiology

Although the precise prevalence of SOV aneurysm is not available, the general population estimate of prevalence is 0.09% (1–5). For etiology, SOV aneurysms are commonly congenital and represent 0.1% to 3.5% of congenital heart defects (6, 7). SOV aneurysm usually originates from the right coronary sinus (70%–90%), followed by the noncoronary sinus (10%–25%) (Fig. 1) and left sinus (<5%) (6). The underlying mechanism for congenital type is incomplete fusion of the distal bulbar septum (primitive bulbis cordis) and truncal ridges (aortopulmonary septum) resulting in fragility at the junction of aortic annulus and the right aortic sinus media and right portion of the noncoronary sinus (Fig. 2). With continuous pressure over time, this morphologic imperfection leads to an increasing bulge, which can eventually rupture (6, 8).

On the other hand, SOV aneurysms in patients with connective tissue diseases like Marfan syndrome and Loey-Dietz syndrome cause annuloaortic ectasia resulting in dilatation of all three SOVs and thereby progressive effacement of the sinotubular junction (3). Acquired causes for SOV aneurysms include atherosclerosis, bacterial endocarditis, syphilis, and tuberculosis (3); autoimmune disease such as Behcet’s disease (9); degenerative conditions such as cystic medial necrosis; traumatic injury and postoperative after surgical repair of a ventricular septal defect (VSD), aortic valvular diseases, and aortic dissection (3, 6) (Figs. 3, 4).

Normal anatomy and physiology of the aortic root and SOV

The aortic root represents the portion of the aorta and left ventricular outflow tract demarcated by the sinotubular junction superiorly and the basal portions of the aortic valve leaflets inferiorly (Fig. 5). Thus aortic root is made of the aortic valve leaflets, the commis-
The three sinuses of the aortic root between the attachments of the aortic valve leaflets inferiorly and the sinotubular junction superiorly. Each sinus denotes an aortic valve cusp and the three sinus nomenclature into right, left, and noncoronary sinus is based on the originating coronary artery from it. The noncoronary sinus is located above the interventricular septum and a portion of the anterior mitral leaflet; the right sinus lies in vicinity to the interventricular septum and the right ventricular parietal bands while the left sinus is proximal to the anter-

or left ventricular free wall and the anterior mitral leaflet (12).

The sinotubular junction, the relatively constricted segment between the aortic root and ascending aorta, is circular and supports the peripheral attachments of the aortic valve leaflets (10).

The aortic valve leaflets hemodynamically separate the aorta from the left ventricle. The nadirs of attachment of the aortic valve leaflets into the wall of the root in a semilunar fashion gives rise to virtual three-dimensional (3D) ring called as aortic annulus (10, 11).

For accurate measurement, normally the SOVs are measured in double short-axis at the aortic root from coronary sinus to its opposite trigon. Another method for measurement of aortic root is named sinus-to-sinus measurement and usually performed from the right coronary sinus to the noncoronary sinus (Fig. 5) (13). A study of 103 patients with ECG-gated multidetector computed tomography (CT) has shown normal SOV end diastole measurement as 3.2±0.6 cm for men and 2.9±0.5 cm for women (14). The study showed that aortic root diameter was associated strongly with body size and less strongly with systolic and diastolic blood pressure and stroke volume in univariate analyses, while the root diameter were shown to vary with age and body surface area in multivariate analysis (11).

Functionally, SOVs play an important role in aortic valve function. They provide a space to prevent blocking of the coronary artery orifices from the open aortic leaflets. Secondly, they favor the development of eddy currents behind the open leaflets which in turn hold the leaflets away from the aortic wall in a position where they will be promptly caught and closed by blood flow during end systole (10).

### Clinical manifestations and complications of SOV aneurysms

SOV aneurysms may manifest at any age given their congenital and acquired etiologies. The clinical manifestations vary with asymptomatic presentation of the incidentally discovered unruptured aneurysms to severe aortic insufficiency and heart failure of the ruptured aneurysms (15).

### Unruptured SOV aneurysms

Overall, unruptured aneurysms are asymptomatic and are incidentally detected during imaging workup of heart murmurs or abnormal cardiomeadiastinal silhouette on radiograph (16). Rarely, they may present with dyspnea, palpitations, arrhythmias, or angina chest pain. Thrombus can form in large SOV aneurysms (Fig. 6) with subsequent risk of catastrophic embolization or endocarditis (15).

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**Main points**

- Sinus of Valsalva (SOV) aneurysm is dilatation of one or more of the aortic sinuses, which is a rare but important aortic root defect that can be a cause of some serious cardiac sequelae.
- SOV aneurysms may manifest at any age given their congenital and acquired etiologies.
- Cross-sectional imaging plays a pivotal role in diagnosis, presurgical planning, and postsurgical follow-up of SOV aneurysms.
- Surgery has been conventionally and commonly used treatment option for repair of SOV aneurysms; either ruptured or unruptured. However, recently, various percutaneous closure techniques are gaining popularity in certain scenarios.

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**Figure 1.** Three-chamber reconstructed view of the heart from ECG-gated cardiac CT angiography data demonstrates unruptured aneurysm from the noncoronary sinus of Valsalva (SOV) (black asterisk) protruding into the left atrium (black arrow).

**Figure 2.** Illustration of the heart and root of aorta showing the pathogenesis for congenital sinus of Valsalva aneurysm.
of systemic embolism and stroke (17). Mass effect from large SOV aneurysm can distort or obstruct the coronary ostia thereby leading to myocardial ischemia and infarction (18–20). Both ruptured and unruptured SOV aneurysms can be commonly complicated with aortic regurgitation (AR) which occurs in 30%–50% of patients; therefore requiring aortic valve function evaluation with echocardiogram or magnetic resonance imaging (MRI) (21) and need for aortic valve replacement at the time of surgical fixation of the aneurysm. Based on the location and extent of SOV aneurysm, mass effect on adjacent cardiac structures can occur with impairment in tricuspid and mitral valves function, or partial obstruction of the right ventricular outflow tract. Few case reports describe their rare complications including aneurysmal dissection into the muscular interventricular septum with resultant arrhythmias, heart block, (22) and infective endocarditis (23).

SOV pseudoaneurysm is extremely rare. It is an outpouching of any of the three SOVs resulting from a deficiency in the tunica media and intima. The vascular lumen is contained by either adventitia only, clotted blood, or surrounding structures. SOV pseudoaneurysm may be spontaneous, traumatic, or infective (24, 25). It can lead to the same complications as the true aneurysm.

Ruptured SOV aneurysms
A ruptured SOV aneurysm is a potentially fatal complication. The size, location, and rapidity of rupture are the major elements predicting clinical consequences. The most common sinus to rupture is right coronary or noncoronary sinuses with the right ventricle being the most common site of rupture (Fig. 7), followed by the right atrium (26, 27). Other less common sites of rupture in descending order include right ventricular outflow tract, left ventricle, the interventricular septum, left atrium, and extracardiac space (26). Rupture into the extracardiac space although rare has generally higher mortality with critical complications of cardiac tamponade (14). Symptoms of rupture include substernal chest pain, abdominal pain, and dyspnea. Not uncommonly, patients may present with acute heart failure, hemodynamic compromise, or sudden cardiac death (20). Ruptured SOV aneurysms can predispose to formation of para-aortic abscess and endocarditis (15).

Multimodality imaging of SOV aneurysms
SOV aneurysms vary in size from subtle dilatation of an aortic sinus to overt crescentic or windsock-shaped exophytic projection.

Figure 3. a–c. Acquired post-traumatic SOV pseudo-aneurysm in a 56-year-old man. Candy cane multiplanar reconstruction early arterial phase (a) and axial late arterial phase (b, c) contrast-enhanced CT chest images show contained rupture of noncoronary SOV with pseudoaneurysm formation into the oblique pericardial sinus (yellow arrow at the site of rupture) with flattening of the left atrium (yellow arrowheads) due to external compression.

Figure 4. a–c. Acquired SOV pseudoaneurysm (white arrow) in a 67-year-old man following aortic valve replacement. Axial (a, b) and oblique sagittal reconstructed (c) images from contrast-enhanced ECG-gated chest CT show contained rupture of left coronary sinus of Valsalva (Lt CS) with pseudoaneurysm formation (c, white asterisk) into the aortomitral space (black asterisk) with mass effect on the left atrium (LA) posteriorly and left main coronary artery (LMCA) anteriorly due to external compression (AA, ascending aorta; LV, left ventricle; RA, right atrium).
from the body or apex of the sinus (3). They can also manifest as saccular outpouching or as mass. Many classification methods have been described for SOV aneurysms, with Sakakibara classification of ruptured aneurysms based on originating cusp and the receiving/drainage chamber being the most commonly used method in the surgical literature (Figs. 8, 9, 10; Table) (28).

Chest radiograph findings are nonspecific and are based on SOV aneurysm location, size, and presence or absence of rupture. Patients can present with abnormal cardiome diastinal silhouette and minimal to moderate increased pulmonary vascularity. However, patients’ right ventricular outflow tract obstruction can manifest with decreased pulmonary vascularity (29).

Conventional angiography, the gold standard test in the past, continues to be used during percutaneous intervention settings (30). However, noninvasive cross-sectional imaging modalities, consisting of echocardiography, cardiac CT, and cardiac MRI have essentially replaced it.

Transthoracic echocardiography is the first screening modality of choice given its good sensitivity, wider availability, and portability (31). Many studies have reported >90% accuracy of echocardiography for detecting the SOV aneurysm with common fallacy being the incorrect detection of the rupture site (23). Transesophageal echocardiography
The effective radiation dose depends on the ECG gating technique: it is 1–6 mSv for prospective triggering (during diastole), which is used if there is regular heart rate <65 bpm; but it is 10–15 mSv for retrospective gating, which is used if there is irregular heart rate, heart rate >65 bpm, or if dynamic information about aneurysm filling and emptying or movement in different phases of the cardiac cycle is required.

If retrospective gating is used, tube current dose modulation is performed since dose modulation is known to reduce the radiation dose.

A 64-detector CT scanner or higher detector scanner is the preferred technique and subsequently images are sent to dedicated 3D workstation for multiplanar and volume rendered technique postprocessing for accurate localization, measurement, and for providing road map to clinicians to decide about the management.

Multiplanar cardiac MRI plays an important role in the SOV aneurysm assessment due to its lack of ionizing radiation, ability to quantify ventricular functions and aortic regurgitant fraction, better temporal resolution, and assessment of wall motion abnormalities. However, MRI has lower spatial resolution as compared with CT. Combination of various MRI sequences such as balanced steady state free precession (SSFP)/bright blood imaging, single-shot turbo spin-echo, black blood imaging (Fig. 12) and contrast-enhanced magnetic resonance angiography allows accurate assessment of the origin and size of SOV aneurysms, thrombosis of aneurysm, and its relation with the surrounding cardiac and mediastinal structures (4, 30).

SOV aneurysms may be associated with several other congenital cardiac abnormalities (4) and cross-sectional imaging can be useful for their assessment. SOV aneurysms have been associated with bicuspid aortic valves (Fig. 13) in approximately 10% of cases (35). Bicuspid aortic valves due to accelerated degeneration predispose to aortic aortopathy manifesting as aortic valvular stenosis (36), ascending aorta and aortic root dilatation out of proportion to hemodynamic factors, and true SOV aneurysms (37–39). The most commonly reported associations are VSDs (30%–60% of patients). Other associated conditions include aortic insufficiency (20%–30%) (35), aortic stenosis, infundibular pulmonary stenosis, left ventricular noncompaction (40, 41), atrial septal defect, coronary...
anomalies (aberrant left coronary artery with separate origins of the left anterior descending and circumflex arteries) (35), left-sided superior vena cava (42), atrial septal defect (43), patent ductus arteriosus, and hypertrophic obstructive cardiomyopathy (44).

**Table. Classification system of congenital SOV aneurysms by Sakakibara and Konno (25)**

<table>
<thead>
<tr>
<th>Types of congenital SOV aneurysm</th>
<th>Description</th>
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<tbody>
<tr>
<td>I</td>
<td>The aneurysm originates in the left portion of the right sinus, protrudes forward and ruptures into the right ventricle near the pulmonary valve. The concurrent presence of VSD under the pulmonary valve is frequent.</td>
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<tr>
<td>II</td>
<td>The aneurysm originates in the mid portion of the right sinus, protrudes and ruptures into the right ventricle. A concurrent VSD is uncommon.</td>
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<tr>
<td>III</td>
<td>The aneurysm originates in the posterior portion of the right coronary sinus. IIIv: The aneurysm projects into the right ventricle behind the septal leaflet of the tricuspid valve after penetrating the membranous septum. IIIa: The aneurysm protrudes into the right atrium. VSD is rarely encountered.</td>
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<tr>
<td>IV</td>
<td>The aneurysm originates in the right portion of the noncoronary sinus and ruptures into the right atrium. A combined VSD is uncommon.</td>
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SOV, sinus of Valsava; VSD, ventricular septal defect.

**Treatment planning and management of SOV aneurysms**

Surgery has been conventionally and commonly used for repair of SOV aneurysms. Following successful surgical repair of a ruptured aneurysm, the prognosis is excellent with 10-year survival rates of 90%–95% (45, 46). Surgical repair is indicated for ruptured aneurysms and unruptured SOV aneurysm with associated congenital defects like VSD, or complications like aortic regurgitation, mitral valve incompetence, right ventricular outflow obstruction, infection, and myocardial ischemia (47). The three surgical routes described consist of aortotomy through the aortic root, through ruptured cardiac chamber site of aneurysm, and combination of both (16, 48).

Recently, various percutaneous closure techniques (45, 46) are gaining popularity in certain scenarios; for example with the rupture opening site less than 9 mm in diameter and the distance between the SOV opening of the ruptured aneurysm and the coronary artery not less than 5 mm (49). Various reported closure devices used in such condition include Rashkind umbrella, septal and ductal occluder devices, and Amplatzer vascular plug (16).

Patients with unruptured, stable, or asymptomatic SOV aneurysms are usually followed clinically and with imaging. Society guidelines regarding management of aortic aneurysm in general are applicable for unruptured SOV aneurysm (48). One study of 53 cases proposed for anticoagulation of patients with unruptured stable aneurysms, and 6-monthly imaging follow-up. In this study, surgical repair was performed in the presence of symptoms, or the affected sinus size more than 50% of the average size of the other two normal sinuses, compressive or distortive effects on surrounding chambers, or with size increase on follow-up imaging (46, 50).

**Conclusion**

SOV aneurysm is a rare important aortic root defect, which can be a cause of some serious cardiac sequel, but easy to remain undiagnosed in busy clinical practice setting. Hence increased awareness of
this entity and prompt accurate diagnosis is important, to prevent fatal event from untreated ruptured SOV aneurysm. Gated multidetector CT and MRI play an important role as noninvasive imaging modalities for their evaluation.

Acknowledgements

We thank Pam Curry, Medical Illustration Department, UT Southwestern Medical Center for her valuable help in supplying the medical illustrations and animations of this work.

Conflict of interest disclosure

The authors declared no conflicts of interest.

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Cross-sectional imaging of sinus of Valsalva aneurysms

Figure 13. a, b. A 59-year-old woman with history of ascending aortic tube graft repair for Type A aortic dissection extending into descending aorta presents with SOV aneurysm. Short axis contrast-enhanced axial (a) and oblique coronal reconstructed (b) CT images through the aortic root demonstrates partially calcified bicuspid aortic valve (white arrow) and SOV aneurysm from noncoronary sinus (black arrow). Note previous Type B aortic dissection in the descending aorta with partial thrombosis of the false lumen.


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