Imaging pitfalls, normal anatomy, and anatomical variants that can simulate disease on cardiac imaging as demonstrated on multidetector computed tomography

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Abstract
Advances in computed tomography have led to continuous improvement in cardiac imaging. Dedicated postprocessing capabilities, faster scan times, and cardiac gating methods reveal details of normal cardiac anatomy and anatomic variants that can mimic pathologic conditions. This article will review normal cardiac anatomy and variants that can mimic disease. Radiologists should be familiar with normal cardiac anatomy and anatomic variants to avoid misinterpretation of normal findings for pathologic processes.

Keywords
Cardiac computed tomography (CT), cardiac CT anatomy, cardiac anatomy, anatomical variants, imaging pitfalls

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Introduction
Substantial advances in multidetector computed tomography (CT) scans, hardware, software, and postprocessing capability have led to marked improvement in the display of cardiac anatomy and pathology. The use of CT scanners with 64 or more detectors is now standard and ECG-gating capabilities are widely available. In this article, we review normal cardiac anatomy and commonly encountered pitfalls using a chamber-specific approach.

Technique
Current CT scanners produce temporal resolution below 250 ms and in some instances less than 100 ms. Although this permits adequate assessment of cardiac structures during conventional CT acquisitions, ECG-gating should be used to optimize imaging and reduce the potential for artifacts that may lead to misinterpretation. The use of beta-blockers to reduce rapid heart rates and coughing for breath-holding are important adjunctive strategies to reduce artifacts.

For most anatomic questions, prospective ECG-gating that provides a snapshot image of the heart during a small part of the cardiac cycle is sufficient and allows radiation sparing. If functional information is required, retrospective ECG-gating acquired throughout the cardiac cycle is necessary. Delayed imaging may also be useful in certain circumstances. Reformatted images are often a critical part of evaluation of cardiac structures. Multiplanar images can be reconstructed in specific cardiac planes to highlight normal anatomy and variants. Maximum intensity projection and volumetric images may provide additional perspective that may be valuable to the interpreting and referring physicians.

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Cardiac anatomy and variants

Right atrium

The right atrium develops from the primitive right atrium and sinus venosus. The sinus venosus forms a smooth wall called the sinus venarum and receives blood from the inferior vena cava, superior vena cava, and coronary sinus. The primitive right atrium forms the trabeculated aspect of the right atrial appendage. The right atrial appendage has a triangular or pyramidal shape with a wide base opening and rough trabeculation of the pectinate muscles (1,2). The pectinate muscles in the RAA may be misinterpreted as a mass or thrombus (Fig. 1). Careful observation of the parallel course of the pectinate musculature to the wall of right atrium on axial or reformatted images will confirm this normal structure (3).

Fig. 1. A 44-year-old man with atrial fibrillation. A pulmonary vein CT for pre-ablation mapping shows multiple small filling defects abutting the lateral wall of the right atrium (arrows), compatible with prominent pectinate muscles. Note contrast opacification in the right heart due to poor timing. LA, left atrium; LV, left ventricle; RA, right atrium; RV, right ventricle.

Fig. 2. A 62-year-old man with chest pain and prior myocardial infarction. Multiplanar images of a coronary CTA for coronary artery evaluation show the normal crista terminalis identified as a smooth soft tissue ridge along the posterolateral wall of right atrium on axial (white arrow in a), coronal and sagittal planes (black arrow in b, c). AO, aorta; RVOT, right ventricular outflow tract; SVC, superior vena cava.
The crista terminalis is a vertically orientated smooth muscle ridge extending from the superior vena cava to the inferior vena cava. It represents the fusion line between the primitive RA and the posterior smooth wall of the sinus venosus portion of the right atrium (1). Some authors believe that there is an internodal conductive pathway between the sinoatrial and atrioventricular nodes within the crista terminalis, which traverses the Eustachian region, and enters the atrioventricular node posteriorly via the coronary sinus (4). Since the size and shape of the crista terminalis can be variable, it may be mistaken for an intracardiac mass or thrombus (3,5–8) (Fig. 2a–c).

The Eustachian valve is located at the junction of the inferior vena cava and right atrium. This valve is associated with a Chiari network, which is a collection of fibrous structures. The Eustachian valve is commonly seen on MRI (2), less often appreciated on CT due to the inflow of unopacified blood from the inferior vena cava (9). The Eustachian valve directs blood flow towards the foramen ovale and normally regresses during embryonic development. It is typically a thin and mobile structure (2) (Fig. 3).

Fig. 3. A 61-year-old woman with head and neck cancer. Chest CT with contrast was performed for restaging. Normal Eustachian valve is seen as a paper-thin structure at the junction of IVC and RA on axial (black arrow in a), coronal (white arrow in b), and sagittal planes (black arrow in c). IVC, inferior vena cava; PA, pulmonary artery.
The Thebesian valve is located within the right atrium at the orifice of the coronary sinus. It prevents reflux of blood from the right atrium into the coronary sinus (10). Both of these valvular structures can be variable in size, shape, and appearance and should not be mistaken for neoplasm, thrombus, or inflammation (11).

**Left atrium**

The left atrium is the most posterosuperior chamber and receives pulmonary venous return. It is attached to the left atrial appendage. The primitive left atrium forms a single pulmonary vein which later divides into four pulmonary veins. The posterior wall of the primitive left atrium, which normally receives the four pulmonary veins, becomes the smooth portion of the left atrium proper. The remainder of the primitive left atrium becomes the trabeculated portion or left atrial appendage (1) (Fig. 5).

Compared to the right atrial appendage, the left atrial appendage is more tubular in shape, has a narrower base, and contains fewer trabeculated pectinate muscles (2). The pectinate muscles can have a linear or globular appearance and can be misinterpreted as an intracardiac mass or thrombus (12). This issue can be resolved by close observation of the course of the pectinate muscles, which are parallel in configuration. In contrast, thrombus is seen as a focal filling defect. If doubt exists, reformatted images will confirm the diagnosis (3).

Blood flow in the left atrial appendage is sluggish and therefore prone to thrombus formation (Fig. 7a, b). The presence of thrombus may alter patient management, particularly if left atrial appendage closure devices are being considered. Dependent contrast and unopacified blood causing a contrast-fluid level in the left atrium can lead to the unopacified component being confused with thrombus (3). Repeat CT with delayed scanning in the venous phase (approximately 30 s later) at the same level will show a well opacified left atrial appendage although echocardiography may be necessary for clarification. Thrombus in the left atrium and left atrial appendage is often associated with atrial fibrillation or cardiac chamber and valvular abnormalities (13).

The left atrial oblique vein is an embryologic remnant of the left superior caval vein. The left superior caval vein is enclosed within the Marshall ligament, which is a fold of the visceral pericardium containing muscle, vascular, and nervous structures (14,15). The ligament of Marshall courses obliquely above the left atrial appendage and lies laterally to the left superior pulmonary vein bundle in the epicardial aspect of the left atrial fold, forming the left posterior crest (16). It is also known as the Coumadin or Warfarin ridge, or Q-tip. This can be mistaken as a pedunculated mass or thrombus (17). Historically, this structure was misinterpreted as a thrombus on echocardiography, resulting in anticoagulation therapy with Warfarin. The Q-tip name derives from an appearance on echocardiography that simulates a Q-tip. This normal structure is well visualized on multiplanar rendering (MPR) imaging (Fig. 8a–c).

The interatrial septum is thin and often not well visualized on imaging studies. Normal fat deposition within the septum measures between 0.9–9.9 mm in thickness (18). Due to advanced age or obesity, increased fat deposition occurs and makes the septum easier to visualize. The most common abnormality of the interatrial septum is lipomatous hypertrophy of the interatrial septum, defined as fatty deposition within the interatrial septum of more than 2 cm in thickness that spares the fossa ovalis (19). This is present in up to 2.2% of patients and is occasionally associated with pulmonary emphysema and atrial arrhythmia (20).

The characteristic features include a smoothly margined, non-enhancing dumbbell-shaped fatty mass sparing the fossa ovalis (18–20) (Fig. 9).

Embryologic development of the interatrial septum includes normal thinning of the septum at the fossa ovalis (Fig. 10) (14,15). This physiologic thinning at the fossa ovalis can simulate an atrial septal defect (21). Lack of flow between the left atrium and the right atrium on dynamic studies in the region of the thinned septum confirms this normal appearance. If the findings are equivocal, echocardiography or MRI can be used for further evaluation.

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**Fig. 4.** A 66-year-old man with chest pain. CTA of the chest performed to rule out pulmonary embolus shows an incompetent Thebesian valve (arrow). CS, coronary sinus.
Seventy-five percent of septum primum and septum secundum tissues fuse after birth. If the foramen ovale (FO) is covered but does not completely seal, it results in a patent foramen ovale (PFO). A PFO is a persistent valvular-like connection between the RA and LA, which usually results in a left to right shunt (Fig. 11). Although most patients with PFO are asymptomatic, they may suffer from desaturated blood and embolic events (22,23).

**Right ventricle**

The right ventricle is located in the most anterior portion of the heart. It has a complex triangular shape and receives blood from the right atrium. It is more trabeculated and thinner than the left ventricle. The unique features of the right ventricle include heavy trabeculation and the presence of the moderator band (2,10).
The right ventricle contains three components: (i) the inlet consisting of the tricuspid valve, chordae tendinae, and papillary muscles; (ii) the trabeculated apical myocardium; and (iii) the infundibulum, or conus, which corresponds to the smooth outflow tract (24).

The landmarks separating the right ventricle inlet and outlet are variably reported to be septomarginal trabeculation (25) or the infundibuloventricular fold (26).

In the embryologic period, the fetal heart forms trabecule carneae, numerous muscle bundles running in the ventricles and locating on the inner ventricular surfaces. During embryonic development, division of cardiomyocytes leads to formation of typical trabeculae (27).

The prominent muscular bands within the right ventricle are composed of the parietal band, septomarginal band, septal band, and moderator band (28,29).

The most commonly seen septomarginal trabeculation is the moderator band, a heavily trabeculated muscle and a unique feature of the right ventricle (Fig. 12). It is a tissue ridge that extends across the right ventricular apex from the anterior papillary muscle to the interventricular septum. It contains the right bundle branch (2,10). The displaced moderator band can simulate a double-chambered right ventricle (30) and a prominent moderator band in right ventricular enlargement or hypertrophy can be mistaken for thrombus or tumor (31).

The right ventricle contains three papillary muscles: the anterior, posterior, and medial papillary muscles. Each muscle attaches to two cusps (2,32). The most commonly seen muscle is the anterior papillary muscle (Fig. 13a–c). If prominent, this structure can be mistaken for an intracardiac mass or thrombus.

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**Fig. 6.** A 42-year-old woman with atrial fibrillation. A pulmonary vein CT for pre-ablation mapping shows a filling defect in the left atrial appendage. Its course on the dynamic images appears to be a pectinate muscle (arrow). RAA, right atrial appendage.

**Fig. 7.** A 79-year-old woman with atrial fibrillation. CT of the pulmonary vein prior to left atrial appendage ligation showed a filling defect in the left atrial appendage (arrow in Fig. 7a) which was filled in on the delayed image (arrow in Fig. 7b). This is sluggish flow in the left atrial appendage simulating left atrial appendage thrombus. LAA, left atrial appendage; MPA, main pulmonary artery.
If doubt persists, reformatted images will confirm the diagnosis (3).

Non-specific fat deposition in the right ventricle is found in up to 17% of asymptomatic subjects and is most often encountered in the elderly. Common locations are the superior wall of the base, the middle segment of the right ventricle, and the right ventricular outflow tract. Fat deposition can mimic arrhythmogenic right ventricular dysplasia or prior myocardial infarction involving the right ventricle (33) (Fig. 14). Functional assessment can be used to differentiate variants from a diseased right ventricle.

Fig. 8. A 62-year-old man with chest pain. Coronary CTA was performed to evaluate coronary artery. On axial (a), and multiplanar reformatted coronal (b) and sagittal (c) planes, the Ligament of Marshall, an embryonic remnant at the junction of left superior pulmonary vein and left atrium is seen as a smooth tissue ridge (black arrow). PV, pulmonary vein; RCA, right coronary artery.

Left ventricle

The left ventricle consists of the inlet, apical, and outlet portions and is separated from right ventricle by the interventricular septum. The septum typically bulges toward the right ventricle due to pressure differences between the chambers (2).

The left ventricle contains two papillary muscles, the anterior and posterior papillary muscles, which are larger than the papillary muscles of the right ventricle. Chordae tendinae attach the papillary muscles to the mitral valve leaflets (2). The posterior papillary muscle
can have a globular shape and can be misinterpreted as
a mass or thrombus (3). Two unique features of the left
ventricle are a smooth wall and shared annulus between
the mitral and aortic valves. Both anterior and pos-
terior papillary muscles are well appreciated on MPR
images (Fig. 15a–c).
In a minority of the population, false tendons or
false chordae occur as normal variants in the left
ventricle. These false chordae are linear structures
attached at both ends to the endomyocardium (34).
They are asymptomatic and usually incidental findings
on echocardiography (31).
The normal wall thickness of the left ventricular
myocardium is 6–11 mm. Focal wall thinning at the
apex with normal cardiac function in the absence of
symptoms or history of myocardial infarction is

Fig. 9. A 64-year-old man with history of lung nodule. A chest
CT performed for re-evaluation of the nodule shows lipomatous
hypertrophy of the interatrial septum (black arrows). The typical
radiographic findings include a dumbbell-shaped fatty mass and
sparring of the fossa ovalis (white arrow).

Fig. 10. A 74-year-old woman with left ventricular thrombus seen on echocardiogram. Cardiac CT was performed to rule out left
ventricular thrombus. The normal thinning of interatrial septum at the fossa ovalis is seen adjacent to the left atrial pouch, corres-
ponding to the unfused septum primum and secundum on four chamber (double white arrows in a) and sagittal oblique views (double
white arrows in b). This can be mistaken for an atrial septal defect.

Fig. 11. A 37-year-old man with atypical chest pain. Coronary
CTA performed to rule out coronary artery disease shows a
small jet of contrast from left atrium to right atrium, compatible
with patent foramen ovale (arrow). This is more conspicuous on
the dynamic study. No coronary artery disease was found.
**Fig. 12.** A 45-year-old man with chest pain. A chest CTA performed to rule out pulmonary embolism shows a prominent moderator band (white arrow) running between the RV free wall and interventricular septum (black arrow).

**Fig. 13.** A 62-year-old man with chest tightness. Coronary CTA was performed to evaluate the coronary arteries. Multiplanar imaging shows an ellipsoid shaped anterior papillary muscle in the right ventricle on axial (a), coronal (b), and sagittal (c) planes (black arrows). MB, moderator band; T, trabeculation.

**Fig. 14.** A 63-year-old woman with atrial fibrillation. A pulmonary vein CT for pre-ablation mapping shows nonspecific fat deposition in the RV free wall (double arrows).
Fig. 15. A 40-year-old man with atypical chest pain. Coronary CTA was performed to evaluate the coronary arteries. Multiplanar imaging with axial and sagittal planes of anterior (a1, a2) and posterior papillary muscle (b1, b2) are shown. These are also seen on the coronal image (c). APM, anterior papillary muscle; IVC, inferior vena cava; IVS, interventricular septum; MB, moderator band; MV, mitral valve; PPM, posterior papillary muscle; TV, tricuspid valve.
considered a normal variant. This normal wall thinning can be mistaken for an old myocardial infarction (Fig. 16). The key to diagnosis is assessment of wall motion during functional evaluation. This can be done if retrospective ECG-gating cardiac CT has been performed or alternatively by echocardiography or cardiac magnetic resonance imaging (MRI) (35).

The interventricular septum consists of a thin membranous portion located immediately beneath the aortic valve and a larger muscular septum (9). The thin membranous septum can simulate a ventricular septal defect (Fig. 17). Lack of visualization of blood flow between two ventricular chambers assures an intact ventricular septum.

**Sinus of Valsalva**

The sinus of Valsalva of the ascending aorta and main pulmonary artery aligns obliquely to the axial plane. An axial CT slice through the sinus of Valsava will cause sinus of Valsalva to appear distorted and can be mistaken for a sinus of Valsava aneurysm. Sagittal or coronal reformats perpendicular to the aortic and pulmonic valve plane will help to delineate this normal anatomy (36) (Fig. 18a–c).

**Pericardial recesses**

Pericardial reflection covering the heart and mediastinum can be variable and confusing when containing fluid. These pericardial recesses can mimic mediastinal masses, mediastinal lymphadenopathy, or even pulmonary nodules.

Mediastinal structures, including the heart and great vessels, are covered by the pericardium. The pericardium consists of two layers, the outer fibrous layer and the inner double-layered serous sac. The inner layer is composed of visceral and parietal layers, which become pericardial recesses. The tissues of origin determine the locations of the pericardial recesses, which can be categorized into transverse sinus, oblique sinus and pulmonary venous recesses (37,38).

The transverse sinus is situated posterior to the ascending aorta and main pulmonary artery above the left atrium. It is divided into the superior aortic, inferior aortic, pulmonic, and postcaval recesses depending on its location (Fig. 19). The superior aortic recess is the most prominent and most variable pericardial recess (Figs. 20 and 21). The oblique sinus is located posterior to the left atrium and anterior to the esophagus. It is separated from the transverse sinus by a double layer of pericardium (Fig. 22). As the pulmonary veins join the left atrium, the pericardium incorporates the veins resulting in right and left pulmonary venous recesses (Fig. 23). These recesses are usually small, however, the right pulmonary venous recess is frequently seen.

A summary of normal structures and anatomical variants that can simulate pathology is shown in Tables 1 and 2.
Fig. 18. A 56-year-old man with atrial fibrillation. A pulmonary vein CT for pre-ablation mapping shows distortion of the Sinus of Valsalva of the aorta which can simulate the sinus of Valsalva aneurysm (arrows in a). Images perpendicular through the aortic valve (black line in b) will clarify this normal anatomy (c).

Fig. 19. An illustration shows pericardial reflection forming the pericardial recesses from the anterior perspective with the heart removed. OS, oblique sinus; PVR, pulmonic vein recess; TS, transverse sinus. This illustration was modified with permission from: Gray’s Anatomy for Students. 2nd edition. Amsterdam: Elsevier, 2010:122–243.

Fig. 20. A 41-year-old woman with shortness of breath. A chest CT shows prominent superior aortic recess surrounding the ascending aorta simulating a periortic hematoma (arrows). As AO, ascending aorta; AZ V, Azygos vein; Des AO, descending aorta.
Fig. 21. A 72-year-old man with shortness of breath. An unenhanced chest CT shows the inferior extension of the periaortic recess (arrows), which may be mistaken as subcarinal lymphadenopathy.

Fig. 22. A 65-year-old man with colon cancer. Restaging chest CT shows prominent oblique sinus filling with fluid (arrows). Note fluid filled in the lumen of the minimally thicken esophagus.

Fig. 23. A 43-year-old man with chest pain. A chest CTA performed to rule out PE shows a small outpouching of fluid from the sleeve of right inferior pulmonary vein (arrows). This is the common location for right pulmonary venous recess.

Table 1. The table summarizes normal structures in the atria that can simulate pathology and how to avoid misinterpretations.

| Cardiac chambers | Normal structures                  | Mimickers          | Remarks                                |
|------------------|-----------------------------------|--------------------|---------------------------------------|
| Right atrium     | Prominent pectinate muscle        | Tumor or thrombus  | Re-evaluation on MPR                  |
|                  | Prominent crista terminalis       | Tumor or thrombus  | Re-evaluation on MPR                  |
|                  | Prominent Eustachian and Thebesian valves | Tumor or thrombus | Re-evaluation on MPR                  |
| Left atrium      | Prominent Ligament of Marshall, Pectinate muscle | Tumor or thrombus | Re-evaluation on MPR                  |
|                  | Sluggish flow in left atrial appendage | Thrombus           | Repeat delayed scan at 30 s or further investigate with echocardiogram |

(continued)
Conclusion
Awareness of normal cardiac anatomy and anatomic variants will assist cardiothoracic imagers in avoiding misinterpretation of normal findings as pathologic processes.

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Conflict of interest
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Table 1. Continued.
| Cardiac chambers | Normal structures | Mimickers | Remarks |
|------------------|------------------|-----------|---------|
| Right ventricle  | Displaced moderator band | Double chamber right ventricle | Further evaluation with echocardiogram or MRI |
|                  | Prominent moderator band | Tumor or thrombus | Re-evaluation on MPR |
|                  | Prominent anterior papillary muscle | Tumor or thrombus | Re-evaluation on MPR |
|                  | Non-specific fat deposition in the RV free wall | Arrhythmogenic Right Ventricular Dysplasia | Further evaluation with echocardiogram or MRI |
| Left ventricle   | Prominent posterior papillary muscle | Tumor or thrombus | Re-evaluation on MPR |
|                  | Normal apical thinning | Prior myocardial infarction | Further evaluation with echocardiogram or MRI |
|                  | Membranous septum | VSD | Further evaluation with echocardiogram or MRI |

Table 2. Normal structures in the ventricles that can mimic diseases and appropriate investigations to prevent misinterpretations.
| Cardiac chambers | Normal structures | Mimickers | Remarks |
|------------------|------------------|-----------|---------|
| Right ventricle  | Displaced moderator band | Double chamber right ventricle | Further evaluation with echocardiogram or MRI |
|                  | Prominent moderator band | Tumor or thrombus | Re-evaluation on MPR |
|                  | Prominent anterior papillary muscle | Tumor or thrombus | Re-evaluation on MPR |
|                  | Non-specific fat deposition in the RV free wall | Arrhythmogenic Right Ventricular Dysplasia | Further evaluation with echocardiogram or MRI |
| Left ventricle   | Prominent posterior papillary muscle | Tumor or thrombus | Re-evaluation on MPR |
|                  | Normal apical thinning | Prior myocardial infarction | Further evaluation with echocardiogram or MRI |
|                  | Membranous septum | VSD | Further evaluation with echocardiogram or MRI |
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