INTRODUCTION

As imaging modalities evolve, the detailed anatomy of the heart, including congenital anomalies, can be detected more easily. However, even with such advancements, anomalies involving the coronary sinus (CS) have received relatively little attention, perhaps because they do not usually cause clinical symptoms or functional disturbances.

However, precise diagnosis of anatomic variants of the CS and its tributaries is crucial to correct the performances of various interventional procedures, such as cardiac resynchronization therapy and percutaneous transvenous mitral annuloplasty (1-3). Additionally, cardiac surgeons must be aware of the anatomy of the CS, as it is used as a route for injecting cardioplegic solutions to minimize cardiac activity and to allow the operation to be performed in a still and bloodless field. This process also allows surgeons to reduce the aortic cross-clamping times and avoid interruptions. Retrograde cardioplegia utilizes the uniform CS venous system, because it has a more uniform venous network and is not affected by coronary artery diseases (3). In patients with severe coronary artery diseases, antegrade cardioplegia may fail to supply the cardioplegic solution to the myocardium sufficiently due to stenotic or occluded arteries. Thus, in such situations, retrograde cardioplegia is preferred. However, failure to recognize congenital anomalies of the CS, such as atresia of the CS orifice or agenesis of the CS, may lead to difficult or failed retrograde coronary venous catheter insertions.

Congenital anomalies involving the CS can be classified into four groups based on the following criteria (4, 5): 1) enlargement of the CS with or without a left-to-right shunt, 2) absence of the CS, 3) atresia of the right atrial ostium of the CS, and 4) hypoplasia of the CS. These anomalies are considered to be the faulty embryologic development results from the heart.

The purpose of this pictorial review is to illustrate and classify CS congenital anomalies, and to present several incidentally founded cases. The cardiac venous system and its embryologic development are also being described in detail to familiarize radiologists with various congenital anomalies of the CS.

Index terms
Coronary Sinus
Congenital Anomaly
Congenital Heart Disease
Multi-Detector Computed Tomography
NORMAL ANATOMY OF THE CARDIAC VENOUS SYSTEM (FIG. 1)

A recent anatomic classification divides the cardiac veins into two main groups: tributaries of the greater cardiac venous system (CVS), and, tributaries of the lesser CVS.

The Greater Cardiac venous System

The greater CVS is divided into two subgroups: CS tributaries and non-CS tributaries. The CS tributaries include the CS as the major component of the CVS and collectively drains about 75% of the coronary venous blood (3). The CS is approximately 45 mm in length and 10-20 mm in diameter. The CS lies along the posterior atrioventricular groove and empties into the right atrium through the orifice guarded by the thebesian valve.

The first branch of the CS is the posterior interventricular vein (or the middle cardiac vein), which courses in the posterior interventricular groove. The next main branches are the posterior vein of the left ventricle and the left marginal vein. The CS continues as the great cardiac vein, which courses in the left atrioventricular groove (6). The anatomic landmark of the junction between the CS and the great cardiac vein junction is the orifice of the oblique vein of Marshall (the oblique vein of the left atrium), the valve of Vieussens, or the left margin of the myocardial sleeve of the CS. On cardiac CT imaging, the valve of Vieussens can be identified as an annular narrowing on the external surface (Fig. 1) (1). Among these tributaries, the left marginal vein and the posterior vein of the left ventricle are typically used for cardiac resynchronization therapy (7), and thus, the variability of these veins should be assessed with imaging prior to therapy.

The Lesser Cardiac venous System

The lesser CVS is made up of the thebesian vessels that form the subendocardial and intramyocardial communicating network. It drains the remaining coronary venous circulations directly into the cardiac chambers (mostly the right atrium and the right ventricle).

EMBRYOLOGIC DEVELOPMENT OF THE CARDIAC VENOUS SYSTEM

Normal venous development is a complex process of progres-
Without Left-to-Right Shunt into the Coronary Sinus

When the CS receives anomalous systemic venous return, it becomes enlarged. Possible anatomic variations in this subgroup include a) persistent left superior vena cava (PLSVC) confluent with the CS, b) partial anomalous hepatic venous return to the CS, and c) continuity of the inferior vena cava with the left super-

**CONGENITAL ANOMALIES OF THE CORONARY SINUS**

Congenital anomalies of the CS were classified into four types by Mantini et al. (5) in 1966 (Table 1, Fig. 3).

**Enlargement of the Coronary Sinus**

Enlargement of the CS may result from many acquired causes, including ventricular systole of the cardiac phase, heart failure, cardiomyopathy, or atrial fibrillation (1). Meanwhile, increased blood flow volume through anomalous congenital communications also causes enlargement of the CS. Such anomalies can be divided into two groups, depending on the existence of a left-to-right shunt into the CS.

**Table 1. Classification of Anomalies Involving the Coronary Sinus**

<table>
<thead>
<tr>
<th>Type</th>
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| 1. Enlargement of the coronary sinus | 1) Without left-to-right shunt into the coronary sinus  
   a) Persistent left superior vena cava  
   b) Partial anomalous hepatic venous connection to coronary sinus  
   c) Continuity of the inferior vena cava with the left superior vena cava through the hemiazygos vein  
  2) With left-to-right shunt into coronary sinus  
   a) Low-pressure shunts  
     - Unroofed coronary sinus  
     - Pulmonary venous connection to coronary sinus  
   b) High-pressure shunts  
     - Coronary artery-coronary sinus fistula  
| 2. Absence of coronary sinus | 1) Atrial septal defect localized to the position normally occupied by the coronary sinus  
  2) Atrial septal defect involving entire lowermost portion of septum  
   a) Persistent common atroventricular canal  
   b) Congenital cardiac disease with asplenia  
| 3. Atresia of the right atrial coronary sinus ostium | 1) Persistent left superior vena cava  
  2) Gross communication of coronary sinus with left atrium  
  3) Multiple communications between coronary sinus and related atria  

Fig. 2. Development of the cardiac veins. Posterior view of the heart of an embryo at four weeks (A), eight weeks (B), and ten weeks (C). By the eighth week, the umbilical veins and the left vitelline vein regress. The left and right anterior cardinal veins connect through a vessel that becomes the left brachiocephalic vein. The right horn of the sinus venosus becomes the posterior wall of the right atrium, and the cardinal veins become the vena cava. The left common cardinal vein is occluded, and the left horn of the sinus venosus forms the coronary sinus and the oblique vein of the left atrium (vein of Marshall). Adapted from Moore KL, Persaud TVN. The developing human: clinically oriented embryology, 7th ed (8).  

Note. LA = left atrium, RA = right atrium
Congenital Anomalies of the Coronary Sinus

Persistent Left Superior Vena Cava

PLSVC (Fig. 3A) is the most common thoracic venous anomaly. The prevalence of PLSVC is reported to be 0.3% in healthy individuals and 4.4% in patients with congenital heart disease (10). This anomaly results from persistent patency of the left anterior cardinal vein, which normally obliterates during development. A PLSVC usually drains into the right atrium through a dilated CS (Figs. 4, 5); however, it may also drain directly into the right atrium in some cases.

This anomaly usually occurs as an isolated lesion, but it can be associated with other congenital cardiac lesions, including the PLSVC draining into the left atrium, interatrial septal defect, or single atrium. The absence of the right superior vena cava also accounts for approximately 20% of PLSVC (Fig. 6) (11). In this case,
both the left and right brachiocephalic veins drain into the PLSVC.

Partial Anomalous Hepatic venous Return to the Coronary Sinus
An anomalous venous channel arising from the liver pierces the diaphragm and the pericardium. It then passes through the posterior aspect of the heart to join the CS (Fig. 3B).

Continuity of the Inferior Vena Cava with the Left Superior Vena Cava through the Hemiazygos Vein
In this condition, instead of connecting to the hepatic segment, the suprarenal segment of the developing inferior vena cava joins the hemiazygos vein, which in turn joins a PLSVC. The PLSVC further drains into the CS, resulting in enlargement (Fig. 3C). This condition may also be associated with other cardiac anomalies, including abnormal positioning of the heart, partial inversion of the abdominal viscera, or polysplenia (5, 9).

With Left-to-Right Shunt into the Coronary Sinus
The CS may also become enlarged when oxygenated blood shunts through anomalous communications. These communications can be divided into two groups: low- and high-pressure, left-to-right shunts.

Communication of the Coronary Sinus with the Left Atrium
This communication may occur either indirectly or directly. In cases of indirect communication, an anomalous vein passes over the lateral wall of the left atrium, and connects the CS with the left atrium. Embryologically, this anomalous vein is consid-

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Fig. 5. Incidentally found persistent left superior vena cava (PLSVC) in a 17-year-old boy who underwent both central venous catheter and chest tube insertion due to a traffic accident.
A. Plain chest radiograph showing the hemodialysis catheter in the right superior vena cava and the subclavian catheter in the PLSVC.
B. Left superior venacavogram confirming that the persistent left superior vena cava drains through the coronary sinus into the right atrium.

Fig. 6. Absence of the right superior vena cava with persistent left superior vena cava (PLSVC).
A. Axial CT image showing the PLSVC (arrow) without the right superior vena cava.
B. The severely dilated coronary sinus (CS) (arrows) drains into the right atrium (RA).
C, D. 3D VR image (C) and curved MPR image (D) showing the PLSVC draining into the severely dilated CS (arrows) inferior to the left atrioventricular groove.
Note.—3D VR = three dimensional volume rendering
Congenital Anomalies of the Coronary Sinus

tered to be a persistent vessel connecting the left atrium or pulmonary vein to the cardinal venous system in response to a partially obstructed CS ostium (12). Blood flow in this channel is expected to be in a left-to-right direction under physiological pressure.

Unroofed CS is a rare anomaly in which communication between the CS and the left atrium is formed directly as a result of the partial or complete absence of the CS roof (Figs. 3D, 7, 8). This entity is the rarest type of atrial septal defect and is often associated with PLSVC (4, 5, 9). This condition usually occurs with a left-to-right shunt; however, if the PLSVC is connected to the CS, systemic venous blood may be diverted to the left atrium via the fenestration. This diversion may result in a right-to-left shunt, which can cause cerebral emboli or brain abscesses.

Pulmonary venous Connection to the Coronary Sinus

This anomaly is either total or subtotal forms, but total anomalous pulmonary venous connection to the CS is the more common form. Under this condition, each of the pulmonary veins forms a pulmonary venous confluence that connects the CS.

Coronary Artery-Coronary Sinus Fistula

Coronary artery-coronary sinus fistula is a rare coronary arte-

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Fig. 7. Incidental unusual communication of the severely dilated great cardiac vein with the left atrium and a partially unroofed coronary sinus (CS) in a 69-year-old female.

A. Axial CT MIP images demonstrate a severely dilated great cardiac vein (white arrow) having a communication with the left atrium just below the left superior pulmonary vein level.
B. The coronary sinus (white arrows) is also dilated and drains into the right atrium.
C. Curved MPR image showing an abnormal communication of the coronary sinus with the left atrium (white arrows). The partially unroofed coronary sinus can be seen (black arrow).

Note.—MIP = maximum intensity projection, MPR = multiplanar reconstruction, RA = right atrium, LA = left atrium

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Fig. 8. Incidentally found partially unroofed coronary sinus (CS) in a 65-year-old woman.

A. Curved MPR image showing a partially unroofed CS, the site of unroofing (arrows), and dense contrast (left-to-right shunt) entering the right atrium (RA).
B. 3D VR image showing a partial direct communication (arrows) between the CS and the left atrium and mild dilatation of the CS.

Note.—MPR = multiplanar reconstruction, 3D VR = three dimensional volume rendering
riovenous fistula. This condition leads to a high-pressure, left-to-right shunt into the CS that is usually dilated. The involved artery becomes elongated and tortuous, and an aneurysm may sometimes form (5).

Absence of the Coronary Sinus
This condition is usually accompanied by other anomalies, such as a PLSVC connected to the left atrium or an atrial septal defect. There are three subdivisions based on the type of atrial septal defect and the existence of additional anomalies (4, 5).

Atrial Septal Defect Localized to the Position Normally Occupied by the Coronary Sinus
This atrial septal defect is located in the posteroinferior angle of the atrial septum that is normally occupied by the CS (Fig. 3E). This condition is considered to arise from the developmental failures of wall formation between the CS and the left atrium. Left-to-right interatrial shunts occur through defect in the left atrial side, which is continuous with the orifice of the CS opening on the right atrial side of the septum. However, when the right atrial pressure exceeds the left atrial pressure, a right-to-left shunt can occur and allows paradoxical emboli to enter the systemic arterial circulation.

Defect Involving the Entire Lowermost Portion of the Atrial Septum
The persistent common atrioventricular canal shows a more extensive malformation of the atrioventricular valves in the form of defects of the atrial septum and the persistent common atrioventricular canal (Fig. 3F). Congenital cardiac disease with asplenia is another syndrome that includes the absence of the CS and splenic agenesis.

Atresia of the Right Atrial Ostium of the Coronary Sinus
In this condition, the CS lies in normal position, but has a blind ostium to the right atrium (Figs. 9, 10). If accompanied by a PLSVC, it carries blood from the CS to the right atrium through the left innominate vein. The condition may occur as an isolated anomaly, although it can be associated with other cardiac malformations (1, 4, 5).

With Functional Persistent Left Superior Vena Cava
Blood from the CS passes the left superior vena cava in a retrograde direction, then moves into the right superior vena cava through the left innominate vein and eventually into the right atrium (Fig. 3G). In this circumstance, the PLSVC is the only outflow channel for the CS. Procedures that interrupt coronary venous drainage, such as ligation of the PLSVC, can lead to myocardial ischemia or coronary venous hypertension (2).

With Gross Communication of Coronary Sinus and Left Atrium
This anomaly has a narrow left superior vena cava joined to the CS with the CS communicating with the left atrium, and is somewhat similar to the unroofed CS mentioned previously (Fig. 3H). The blood flows from the CS through this opening into the left atrium.

With Multiple Communications between Coronary Sinus and Related Atria
In this subtype, there is no left superior vena cava, and blood
Congenital Anomalies of the Coronary Sinus

from the CS flows directly into the related atria (Fig. 3I) (4, 5).

Hypoplasia of the Coronary Sinus

In this rare subtype, some cardiac veins may not join the CS and may individually empty the atrial chambers through dilated thebesian channels due to a failure when joining the coronary sinus (1, 5). Unlike atresia or stenosis of the CS orifice (Fig. 11) which is a focally involved lesion at the CS orifice; hypoplasia and agenesis involve lesions throughout the CS diffusions.

SUMMARY

Congenital anomalies involving the CS can be classified into four groups and several subgroups depending on the presence of additional anomalies. These defects do not usually cause significant symptoms or dysfunctions; however, during interventional treatment for heart disease or retrograde cardioplegia during cardiac surgery, knowledge about the anatomy and the potential anomalies of the CS is important. Recognizing variations of congenital CS anomalies may help avoid errors during these procedures.

REFERENCES

관상정맥동의 선천적 이상: 임상화보

이정은 · 권세환 · 오주형

관상정맥동 기형은 임상 증세와 심장 기능 이상이 드물기 때문에 상대적으로 관심이 적었다. 그러나, 최근 영상 검사 방법의 발달로 심장의 자세한 형태는 물론 관상정맥동 기형을 자세히 연구할 수 있게 되었다. 본 임상화보의 목적은 심장 정맥계의 발생학과 관상정맥동 기형을 분류하고 그 형태와 다양한 증례들을 보여주고자 하는 것이다.

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