Structural Features of the Sinus of Valsalva and the Proximal Portion of the Coronary Arteries: Their Relevance to Retrograde Aortocoronary Dissection

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Introduction and objectives. Retrograde aortocoronary dissection is an unusual complication of coronary angioplasty. Our study provides new structural details of the aortic sinuses and the proximal portions of the coronary arteries, which enable better understanding of several clinical features associated with this complication.

Methods. We studied eight aortic sinus specimens from patients with structural ischemic heart disease using dissection, histologic analysis, and scanning electron microscopy, and compared findings with those in eight control specimens.

Results. We observed the following features: a) in 10 specimens (71%), the left coronary artery diameter was greater than the right; b) the angle that the ascending aorta made with the left coronary artery was acute, whereas that with the right coronary artery was closer to a right angle, thereby possibly providing a better approach for catheterization; c) in contrast to those of the right coronary artery, the periostial wall and sinotubular junction of the left coronary artery were formed by more smooth muscle cells and by a dense matrix of collagen type-I fibers; and d) the aortic sinuses and coronary arteries in structural ischemic heart disease specimens displayed structural alterations that affected the aortic tunica media and the collagen distribution at the sinotubular junction.

Conclusions. The morphological and structural differences observed between right and left sides suggest that the left aortic sinus is more resistant to traction and is, therefore, less prone to iatrogenic dissection. Structural ischemic heart disease is a risk factor that increases the likelihood of aortocoronary dissection.

Key words: Aortocoronary dissection. Aortic sinuses. Ischemic heart disease. PTA.

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Características estructurales de los senos de Valsalva y porción proximal de las arterias coronarias. Su relevancia durante la disección retrógrada aortocoronaria

Introducción y objetivos. La disección retrógrada aortocoronaria es una complicación poco frecuente que sigue a la angioplastia coronaria. Nuestro estudio proporciona nuevos detalles estructurales de los senos aórticos y la porción proximal de las arterias coronarias para un mejor entendimiento de aspectos relacionados con esta complicación.

Métodos. Hemos analizado mediante técnicas de diseción, histología y de microscopía electrónica de barrido los senos aórticos de 8 especímenes con cardiopatía isquémica estructural y se comparan con 8 especímenes control.

Resultados. Hemos observado las siguientes características: a) el diámetro de la arteria coronaria izquierda en 10 especímenes (71%) fue mayor que el de la derecha; b) el ángulo que forma la coronaria izquierda con la aorta ascendente es agudo, mientras que el de la coronaria derecha tiende a ser recto, por lo que puede presentar un mejor abordaje para la cateterización; c) la pared perioscial y la unión sinotubular de la coronaria izquierda contiene, a diferencia de la derecha, más células musculares lisas y una densa matriz de colágeno tipo I, y d) los senos aórticos y coronarios de los especímenes con cardiopatía estructural presentan alteraciones estructurales que alteran la túnica media aórtica y la distribución del colágeno de la unión sinotubular.

Conclusiones. Las diferencias morfológicas y estructurales observadas indican que el seno aórtico izquierdo es más resistente a las tracciones y, por lo tanto, menos propenso a la disección iatrogénica. La cardiopatía isquémica estructural es un factor de riesgo que incrementa la predisposición a la disección aortocoronaria.

Palabras clave: Disección aortocoronaria. Senos aórticos. Enfermedad isquémica. ACTP.
INTRODUCTION

Aortic dissection is an unusual complication of percutaneous transluminal coronary angioplasty (PTCA).\(^1\) Most cases described in the literature occurred after interventions addressing the right coronary artery,\(^1\) and, very rarely, after left coronary artery catheterization.\(^2\) This complication is potentially serious and can lead to acute myocardial infarction or sudden cardiac death.

Using dissection and histological sections, we studied the course and morphological and structural characteristics of the coronary sinuses and the proximal part of the left and right coronary arteries in post-mortem specimens with and without structural ischemic heart disease, with the aim of studying the mechanisms and factors that can make the left coronary artery less prone to retrograde dissection than the right during PTCA.

METHODS

We studied 16 post-mortem hearts that had been previously fixed by immersing them in 10% buffered neutral formalin, while avoiding doing this under pressure via the coronary arteries as this could have distended them and led to the samples becoming distorted. The causes of death were associated with: road traffic accident (n=6), cirrhosis of the liver (n=2), suicide (n=3), cerebral hemorrhage (n=3), and pulmonary thromboembolism (n=2). In total, 8 of the 16 specimens presented structural ischemic heart disease with stenosis of the right and left coronary arteries due to atherosclerosis. There were 10 male and 6 female patients, 30-78 years old (mean ± standard deviation, 55±9 years). The weight of the hearts ranged between 332 and 450 g (380±22 g); the diameters of the left and right coronary artery ostia were 3.7±0.3 cm (range, 2.6-4.2 cm). The right coronary artery ostium (3.7±0.5 mm) was greater (P<.001). Using dissection and optical microscopy techniques, all the specimens showed that the ostium and the first 2-3 mm of the coronary arteries were located within the wall of the ascending aorta or the aortic wall. From here, the initial extraaortic part resected and processed to create histological sections. Two blocks were made from each heart, approximately 7 mm thick, which were dehydrated in graded alcohol, embedded in paraffin, and sectioned consecutively at 10 µm in the frontal plane. The sections were stained at 60 µm intervals using Masson trichrome and picrosirius red F3BA (Gurr, United Kingdom) protocols at 1% dilution. Using a polarized light microscope, the collagen fibers stained with picrosirius red present birefringence, indicating the presence of submicroscopic units oriented along the fiber axis. These subunits are made up of type I and II collagen. Type I is strongly birefringent, with colors ranging from yellow to red; on the other hand, type III collagen is less refringent and appears green.

The structures of the histological sections were measured using image analysis software (SigmaScanPro 5.0, Jandel Scientific, San Rafael, CA, USA). To aid in visualizing the coronary ostia architecture and the ascending aortic wall, we used a scanning electron microscope (Jeol JSM 5600) to examine the 25-30 µm histological sections that had previously been deparaffinized in xylol for 30 min, then afterwards dried at room temperature for 1 h, and covered with gold (BAL-ECT SDC 005 Sputter coater) for 4 min.

Statistical Analysis

The results are expressed as mean ± standard deviation (SD). Statistical analysis was done using the Student t test for independent samples in the case of quantitative variables. P values <.05 were considered statistically significant.

RESULTS

Morphology of the Aortocoronary Junction and Its Proximal Tract

All the hearts studied had 3 aortic leaflets and the coronary arteries originated in the corresponding sinuses of Valsalva (Figure 1). Aortic sinus diameters were 3.7±0.3 cm (range, 2.6-4.2 cm). The right coronary artery ostium (12 hearts, 75%) and the left (12 hearts, 85%) were located below the sinotubular junction. There were significant differences between the diameters of the left and right coronary artery ostia. The diameter of the left coronary artery ostium in 10 specimens (71%) was greater (4.5±0.4 mm) than the right coronary artery ostium (3.7±0.5 mm) (P<.001). Using dissection and optical microscopy techniques, all the specimens showed that the ostium and the first 2-3 mm of the coronary arteries were located within the wall of the ascending aorta or the aortic wall. From here, the initial extraaortic part

ABREVIATIONS

PTCA: percutaneous transluminal coronary angioplasty.
(approximately 2 cm) of the left coronary artery descends parallel to the ascending aorta and forms, with the aortic sinus wall, an angle of 35.5°±11.5° (range, 20°-55°), to course between the pulmonary trunk and the left atrial appendage (Figure 1). In contrast, the initial part of the right coronary artery originates almost perpendicular to the aortic sinus wall forming an angle of 71.5°±8.5° (range, 60°-88°), and passes in front of and slightly to the right of the right atrium, lying between this and the trunk of the pulmonary artery (Figure 1). As it approaches the atrioventricular groove, the right coronary artery descends almost vertically.

### Histological and Structural Study of the Coronary Artery Ostia and the Aortic Wall

Using conventional, polarized light and scanning electron microscopy, histological examination showed that the walls of the sinuses of Valsalva are basically made up of type I collagen in their lower part proximal to where the aortic leaflets attach, where muscle fibers insert into the left ventricle (Figure 2); however, the number of type I collagen fibers decrease as the elastic fibers in the ascending part of the aortic sinuses increase (Figure 2). The aortic wall thickness was 1.8±0.3 mm (range, 1.1-2.4 mm) in the medial portion of each sinus, both in the normal hearts and in those presenting structural ischemic cardiopathy. However, the sinuses where structural ischemic cardiopathy was found presented non-uniform variations in thickness of the elastic lamina of the medial layer and atherosclerotic plaque, at times with hemorrhagic clotting, at the base of the leaflet attachment below its arterial wall (Figure 2).

The upper limit of each sinus at the peak of the line of the semicircular edge of each leaflet is known anatomically as the supravalvular ridge, marking the junction between the sinuses and the tubular part of the aorta. The ridge at the sinotubular junction is mainly made up of elastic and collagenous fibers mixed with smooth muscle cells and fibroblasts. The ridge in the left coronary sinus contains a greater number of smooth muscle cells within a dense extracellular matrix of type I collagenous fibers (Figure 3). In contrast, the right coronary artery has a smaller amount of smooth muscle fibers, which are basically set within type III collagen (Figure 3). The aortic wall thickness at the ridge is 4.3±0.5 mm (range, 3.6-5.1 mm), with significant differences...
between the thickness of each sinus of Valsalva wall and the supravalvular ridge in both coronary arteries ($P<0.001$).

The periostial aortic wall in the sinotubular ridge is characterized by having a prominent tunica media between the internal elastic lamina and the adventitia. This media is predominantly made up of layers of elastic material that alternate with bundles of smooth muscle cells with differing spatial orientation and type I and III collagen fibers (Figure 3). The thickness of the aortic tunica media in this location is very irregular, the longitudinal fibers being mixed with oblique ones. The presence of atherosclerotic plaque and intramural hematoma in the sinotubular ridge produces a thinning of the aortic tunica media (Figure 3), less than 1 mm thick, and its visualization via polarized light shows a non-homogeneous distribution of type I collagen within the sinotubular ridge, like layers of an onion, which decreases on the periostial aortic wall (Figure 3). In one case of progressive atherosclerosis, as found in the 72 year old specimen (Figure 4), the aortic tunica media was characterized by an absence of smooth muscle fibers, immediately above the sinotubular ridge, such that the media was made up of elastic fibers only in this region (Figure 4).

In addition to aortic disease, atherosclerotic plaque affects the most proximal segment of the coronary arteries and is accompanied by marked atrophy of the tunica media with a reduction in elastic and smooth muscle fibers, and at times ulceration, i.e., rupture of the plaque coating due to an increase in pressure promoting thrombosis and coronary obstruction. Such obstruction shows positive staining under polarized light, basically for interstitial type I collagen in the adventitia and media (Figure 4). Finally, it is worth considering the possibility that the plaque is sclerosed and calcified (Figure 4), and the percentage reduction of the lumen is relevant regarding its functional impact.

**Figure 2.** a-d: frontal histological sections stained with Masson trichrome (a, b, and c) and picrosirius red (d) through the aortic sinuses of the right coronary artery (a, c) and left (b, d). It can be seen that the right and left aortic sinuses have more connective tissue fibers (arrows) in the area where the aortic leaflets attach. The pathological leaflets show atherosclerotic plaque (arrowheads) below the arterial wall of the leaflet. In figure d, under polarized light, the difference in staining can be seen between type I collagen fibers (red-yellow) and those of type III collagen (green).

LCA indicates left coronary artery; RAS, right aortic sinus; LAS, left aortic sinus.
The most external layer of the aortic coronary wall is the so-called tunica adventitia, which consists of a network of fibers, basically type I collagen, elastic fibers, adipocytes and macrophages (Figures 1 and 2). No visible alterations in this layer were found in the atherosclerotic arteries studied. The vasa vasorum is normally located in the adventitia, where nerve bundles are also found. The thickness of the tunica adventitia in the aortic wall is 1.2±0.4 mm (range, 0.5-1.8 mm). The aortic tunica adventitia is continuous with the adventitia of both coronary arteries.

**DISCUSSION**

Although the risk of retrogressive dissection in the ascending aorta during PTCA is rare, and that this technique is currently very common and the number of cases has increased, the number of times this serious complication occurs continues to be low. The incidence (0.029%) in our hospital is similar to that in other hospitals, ranging from 0.02 to 0.15%, with an average of 0.059%.

Figure 3. Frontal histological sections stained with Masson trichrome (a, d, f), picrosirius red under polarized light (b, and g) and electron scanning (c) of the sinotubular ridge and left periostial aortic wall (ac) and right (d-g). Note that the sinotubular ridge of the left coronary artery (a, b) has a greater number of smooth muscle cells and type I collagen fibers under polarized light (red-yellow in b) than the right coronary artery (d, e). Under scanning electron microscopy, we can see that the smooth muscle cells (SMC) in the left sinotubular ridge (c) overlap within a dense extracellular matrix, strengthening this part of the coronary ostium. In (f) atherosclerotic plaque (AP) can be seen in the sinotubular ridge of a right coronary artery that affects the tunica media, causing thinning of the aortic wall. In (g) atherosclerotic plaque (AP) appears like layers of an onion under polarized light. Adv indicates tunica adventitia.

Figure 4. Frontal histological sections stained with Masson trichrome (a, b, c) of the sinotubular ridge and periostial aortic wall of the right coronary artery in patients with structural ischemic heart disease. Note in (a) the loss of smooth muscle fibers (arrow) on the aortic wall above the sinotubular ridge. In (b and c) changes can be seen in the aortic tunica media (arrow) which is continuous with the coronary artery tunica media, due to atherosclerotic plaque (AP) and coronary obstruction. In (d and e) cross-sections are shown of the proximal part of an almost obstructed right coronary artery with picrosirius red (d) and polarized light (e). f and g: cross-sections of the proximal part of atherosclerotic plaque (AP) calcified in the left coronary artery with scanning electron microscopy (f) and polarized light (g). Note the atrophy of the arterial tunica media (arrows). L indicates vessel lumen.
Atherosclerotic disease due to myocardial ischemia, which could be a risk factor for increased predisposition to aortocoronary dissection; however, these differences were not found when comparing the atherosclerotic right coronary artery with the left, which indicates that the pathogenesis of the atherosclerotic plaque should be thought of as a set of noxious conditions able to cause endothelial damage, regardless of whether the aortocoronary junction is the right or left although, as stated by Zamir and Sinclair,\textsuperscript{67} it is the aortocoronary junction which is compromised more often. Although one specimen varies from another, the atherosclerotic lesion penetrates the internal elastic lamina and not only affects the distribution of the sinotubular union type I collagen, but there is also thinning of the smooth muscle fibers of the aortic tunica media. In more severe cases, the weakness of the wall occurs in a 1 or 2 mm longitudinal section in the tunica media, where the smooth muscle fibers are replaced by elastic fibers. Aortic weakness in atherosclerotic disease could be a preexisting factor that may play a role in iatrogenic dissection in the face of aggressive interventions such as mechanical traction or contrast-agent injection, as done during PTCA. Another potential risk factor is the presence of total or partial coronary occlusion in its proximal part, as found in some of the specimens, which possibly plays a role radically different to that of degeneration of the aortic tunica media, since percutaneous recanalization is much more complex and requires more aggressive maneuvers than other types of stenosis.

CONCLUSIONS

Our study demonstrates structural differences between the aortic sinuses and the proximal part of the right and left coronary arteries. These differences indicate that the left aortic sinus is more resistant to traction and mechanical pressure than the right and, thus, is less prone to iatrogenic dissection. Atherosclerotic lesions that compromise the aortocoronary junction are a risk factor that increases predisposition to dissection and should be taken into account during PTCA.

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