VARIATIONS IN THE NUMBER AND MORPHOLOGY OF CUSPS OF AORTIC VALVE- ANATOMICAL AND SURGICAL RELEVANCE

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ABSTRACT

BACKGROUND
The human aortic valve is a vital structure and knowledge about its normal anatomy is important in detecting valve diseases and in valve replacement surgeries. The normal aortic valve has three semilunar leaflets or cusps, supported within the three aortic sinuses of Valsalva. The valve is known to exhibit anatomical variations with respect to number, shape and size of cusps, circumference and presence of fenestrations. The aim of the study is to detect and document the variations with respect to the number and morphology of the cusps of aortic valve. A prospective observational type of study on 210 random autopsy cases during a period of 6 months with age ranging from 2½ years to 89 years was conducted in the Dept. of Forensic Medicine, Govt. Medical College, Kozhikode, after getting consent from Institutional Ethics Committee.

MATERIALS AND METHODS
Aortic valves from hearts dissected at autopsy were washed thoroughly and fixed in 10% formalin. Each specimen was numbered systematically. Morphological changes and anatomical variations of the aortic cusps were noted. Relation to cause of death, if any, was also noted.

RESULTS
1. Of the 210 cases studied, 208 aortic valves had three cusps (99.06%), 1 case had a bicuspid aortic valve (0.47%) and one had a quadricuspid valve (0.47%). 2. Fenestrations were seen in 110 valves (52%). 3. Cusps were asymmetrical in 54 aortic valves (25.71%). 4. Lamb's excrescences were seen in 21 cases (10%). 5. Only one case of bicuspid aortic valve with aortic stenosis was associated with sudden cardiac death. Other anomalies were not associated with cardiac cause of death.

CONCLUSION
The anatomical variations and morphological changes occurring in the aortic valve have significant clinical relevance in the diagnosis, prognosis, management and prevention of valve diseases, valve repair and valve replacement surgeries.

KEYWORDS
Aortic Valve, Aortic Valve Abnormalities.


BACKGROUND
An increase in the frequent use of conservative surgical techniques for repairing or replacing damaged valves has generated an interest in the anatomy of valves. Thromboembolism, ruptures, restenosis and calcifications in the prosthetic valves are the major causes of morbidity and mortality in patients. This shows the necessity to develop a substitute of the natural valve, with minimum complications.

The aortic and pulmonary valves are the semilunar valves of the heart. The aortic valve is situated at the end of the aortic vestibule (the smooth left ventricular outflow tract) It has 3 semilunar leaflets or cusps supported within the three aortic sinuses of Valsalva. The free margin of each cusp contains a central fibrous nodule; from each side of this nodule, a thin smooth margin called the lunule extends up to the base of the cusp.

Opposite the cusps, the vessel walls are slightly dilated to form the aortic sinuses. The coronary arteries arise from the anterior and left posterior aortic sinuses. The anterior aortic sinus is also called the right coronary sinus because of the origin of right coronary artery (RCA). The left posterior sinus is called the left coronary sinus because the left coronary artery (LCA) usually arises here. The right posterior sinus is often called the non-coronary sinus, because no coronary arteries arise from it.

Sinotubular junction (STJ) is the point in the ascending aorta, where the aortic sinuses end and the aorta becomes a tubular structure.

Development
The pulmonary and aortic valves are derived from endocardial cushions that are formed at the junction of the truncus arteriosus and the conus. Two cushions, right and left, appear in the wall of the conus. They grow and fuse with each other with the separation of the aortic and pulmonary orifices. The right and left cushions are subdivided into two parts, one part going to each orifice. Simultaneously two more endocardial cushions develop – anterior and posterior. As a result, the aortic and pulmonary openings each have three cushions, from which 3 cusps of the valves develop.

The pulmonary valve is at first ventral to the aortic valve. Subsequently, there is a rotation, so that the pulmonary valve

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comes to the ventral and to the left of the aortic valve. After this rotation, the cusps acquire their definitive relationships as follows. Aortic valve–1 anterior (Right) and two posterior (Left posterior, right posterior); Pulmonary valve–1 posterior, 2 anterior.

MATERIALS AND METHODS
The prospective observational study was conducted in the mortuary, Department of Forensic Medicine, Govt. Medical College, Kozhikode, Kerala. The hearts were obtained from 210 random autopsies. The age of the subjects ranged from 2½ years to 89 years. All cases except those showing decomposition, trauma involving the valve and unknown identity were included in the study.

The hearts, after autopsy, were washed thoroughly in tap water. Weight of the heart and the thickness of ventricles were noted. All the 4 valves–aortic, pulmonary, tricuspid and mitral–were examined. The aortic valves were removed by trimming the ascending aorta 1-2 cm above and below the valves, numbered systematically and immersed in 10% formalin for 24-72 hours.

The valves were stretched, pinned to a wooden board with drawing pins and circumference was measured using a ruler to the nearest millimeter. The horizontal (Intercommissural) length and vertical length of each cusp was also measured. Other anatomical features like asymmetry of the cusps, number of cusps, calcification, presence or absence of fenestrations, other cardiac anomalies were also noted.

Photographs were taken. Other parameters like age, sex, height, weight, associated congenital anomalies, if any, were also recorded.

RESULTS
Number of Cusps
Normal aortic valve with three cusps or leaflets was present in 208 cases out of 210 (99.06%). Quadricuspid aortic valve was seen in one heart (0.47%) (Fig. 1) and bicuspid in one (0.47%) (Fig. 2).

Quadricuspid Aortic Valve
This was found in a 45-year-old male. The cause of death was non-cardiac. The heart weighed 400 gms. Thickness of ventricles was normal. The circumference of the aortic valve was 7 cm. The dimensions of the cusps are shown in Table 1.

From the table, it is evident that one cusp is large, two intermediate and one small. This type comes under the ‘D’ type of Hurwitz and Roberts classification of quadricuspid aortic valves.

All the four cusps had small fenestrations. There were no cardiac anomalies.

Bicuspid Aortic Valve
A 32-year-old male collapsed while playing football and was brought dead to the casualty. Autopsy revealed features of left ventricular failure. The aortic valve had two calcified, distorted cusps, located anteroposteriorly. Circumference of the valve was 3.6 cm. When the two cusps were approximated, there was a large gap. The deceased might have had severe aortic stenosis and regurgitation, leading to left ventricular hypertrophy. The situation was worsened by vigorous exercise.

No Other Cardiac Anomalies were detected

Asymmetry of Cusps
In the present study, 54 aortic valves (25%) showed asymmetrical cusps (Fig. 3) as shown in Table 2.

The intercommissural distance of each cusp was measured in the 21 valves which had 3 cusps of different sizes. The non-coronary cusp was largest in 12, the right coronary in 6 and left coronary in 3 valves

Fenestrations
Out of 210 valves, 110 showed fenestrations (52%) (fig. 4). The size and shape of the fenestrations varied, starting from pinhole size to 0.5 cm. Lowest age in which the valve showed fenestrations was 2½ years– only one cusp had a small hole. But in an 89-year-old male, none of the cusps had fenestrations.

The details of fenestrations and cusps affected is summarized in Table 3. When one cusp alone showed fenestrations, the left cusp was the most affected and non-coronary was the least affected.

Lambl’s Excrescences
Small filiform processes–the Lambl’s excrescences (Fig. 5) were found in 21 valves out of 210 (10%), of which 20 were in adult males and one in an adult female. All three cusps showed Lambl’s excrescences in 3 cases; two cusps in 8 cases and only one cusp showed these processes in 10 cases (5 non-coronary, 4 left coronary and 1 right coronary cusp).

Cause of death in all the 21 cases was non-cardiac and none of the deceased had cardioembolism.

In the present study, the lowest age in which excrescences were seen was 27 years and highest, 88 years.
DISCUSSION

Variations in the Number of CUSPS

The Quadricuspid Aortic Valve (QAV)

The QAV is a rare congenital heart disease. In 1923, Simonds found 2 cases of QAV in 25,666 autopsies, an incidence of 0.008%. After the advent of echocardiography, QAV have been diagnosed more frequently. Feldman et al at the Mayo Clinic detected 8 cases of QAV among 60,446 patients, during echocardiography (Incidence of 0.013%) for the period of 1982-88. For the period 1987-88, 6 cases were identified by the authors among 13,805 examinations (Incidence of 0.043%).

In 1883, Dilg found 2 cases of QAV in a literature review. According to this review, a 34-year-old woman with QAV was reported by Balington in 1862, probably the 1st case of QAV reported. Oktay Tutarel of Hannover Medical School, Germany did a literature search and a total of 186 cases with a QAV was identified.

In 1973, Hurwitz and Roberts classified the QAV according to their anatomic variation. They described 7 variations.

A - Four equal cusps.
B - Three equal and one smaller cusp.
C - 2 equal larger and 2 equal smaller.
D - One large, 2 intermediate and 1 small.
E - Three equal cusps and one larger cusp.
F - Two equal larger and two unequal smaller.
G - Four unequal cusps.

The QAV is said to be associated with other congenital cardiac malformations such as hypertrophic cardiomyopathy or ASD. Coronary arterial anomalies were the most common defect associated with QAV, which was of high significance during aortic valve surgery.

M. Serdar Kuchukoglu et al reported a case of QAV associated with AS and AR in a 26-year-old woman, diagnosed by Two-dimensional echocardiography. 4 cusps were of equal size and of normal texture-Type A.

Jonathen Timperley et al diagnosed 3 cases of quadricuspid aortic valve during echocardiographic examination and one found unexpectedly during aortic valve replacement.
Olson et al. observed an incidence of 1% of QAV on review of 225 patients undergoing surgery for pure aortic insufficiency.

**Embryology**

The embryology of QAV is unknown. A variety of mechanisms have been postulated.

1. Anomalous septation of the truncus arteriosus.
2. Excavation of one of the endocardial cushions.
3. Septation of a normal valve cushion because of inflammation.
4. Division of one of the 3 endocardial cushions that normally give rise to three cusps.

Even though Type A and B were said to be the commonest in literature, in the present study, we found 'D' type of QAV. All the four cusps had small fenestrations. They were too small to produce aortic regurgitation. There were no cardiac anomalies. Origin and branching pattern of coronary arteries were normal.

There is one case report by Clothrap et al. on the presence of large fenestrations in all leaflets of a quadricuspid aortic valve leading to AR.

**Bicuspid Aortic Valve (BAV)**

BAV is the commonest congenital cardiac abnormality with an estimated prevalence of 1-2%. Sir William Osler was one of the first to recognize the bicuspid aortic valve. It is almost three times more common in males than females. A BAV is composed of two leaflets, the commonest configuration of the bicuspid valve has the two commissures located in an anteroposterior direction with left and right cusps.

There can be several associated non valvular lesions. Most patients with BAV have a left dominant coronary circulation. This left coronary can arise from the pulmonary artery. BAV can be associated with other left sided obstructive lesions such as aortic stenosis, coarctation of aorta and ventricular septal defect, suggesting a common developmental mechanism. Specific gene mutation have been isolated.

Linda Cripe, Gregor Andelfinger et al statistically tested whether BAV is associated with genetic inheritance and found out that the heritability of BAV and BAV and/or CVMs were 89% and 75% respectively.

It is now accepted that BAV is heritable. There is a 10% chance of 1st degree relative having a bicuspid aortic valve in patients with BAV disease. Mutations in a gene called NOTCH1, were noted in two families with BAV. So, the American College of Cardiology/American Heart Association recommends that all patients with a 1st degree relative with BAV should be evaluated for BAV and aortopathy.

**Clinical Progression of BAV**

Range from severe aortic stenosis in childhood to asymptomatic disease until old age. In 75%, progressive fibrocalcific stenosis can occur which can lead to sudden death. Paul WM Fedak, Subodh Verma et al have described the Clinical and Pathophysiological Implications of a BAV. Patients with poor lipid profiles and those who smoke are at a high risk of developing aortic stenosis. So, by avoiding these risk factors, chances of developing AS can be minimised.

In the present study, the aortic valve had two calcified, distorted cusps, located anteroposteriorly. There was evidence of aortic stenosis, regurgitation, left ventricular hypertrophy and failure, which led to sudden death.

**Asymmetry of Cusps**

Lee Joseph et al. in a case control study, reported that the presence of asymmetry among AV cusps is strongly associated with the future development of AS. In the present study, 54 aortic valves (25%) showed asymmetrical cusps. A comparison of the present study with that of Lee Joseph et al is shown in Table 4.

In the present study, the number of valves with three equal sized cusps are less than that of Lee Joseph et al and number of valves with three different sized cusps is higher.

A comparison of inter commissural distance of the three cusps is shown in Table 5. According to a study conducted by Marcelo Bisciglì et al., the non-coronary cusp is the largest. In the present study also, non-coronary cusps were the largest when the cusps showed asymmetry. Left coronary cusps were found to be the smallest in the present study as well as in the study conducted by Vollebergh and Becke.

**Fenestrations in Cusps**

Fenestrations of the aortic valve rarely produce significant valvular regurgitation. But, when unusually large or multiple, they can lead to massive aortic regurgitation (AR), mostly in patients with chronic hypertension and/or aortic annular dilation. Rheumatic aortic regurgitation has been reported in a patient with large congenital fenestrations in all three leaflets by Ahmedrezazodati et al.

Hope and Rokitansky were the first to describe aortic cusp fenestrations. Foxe conducted the first systematic study of fenestration of the semilunar valves of the human heart in 300 autopsies and found one or more fenestrations in 82% of all cases. He also concluded that the frequency of fenestrations increases with age.

Acute rupture of the fenestrated cusp, either spontaneously or because of infection can also lead to massive AR resulting in acute left heart failure.

In the present study, out of 210 valves, 110 showed fenestrations (52%). Foxe has reported a higher incidence of fenestrations (82%) The size and shape of the fenestrations varied, starting from pinhole size to 0.5 cm. Lowest age in which the valve showed fenestrations was 2½ year–only one cusp had a small hole. But in an 89-year-old male, none of the cusps had fenestrations. A relation between age and sex with occurrence of fenestrations was not seen in the present study. All the four cusps of the quadricuspid aortic valve showed small fenestrations. A similar finding was also reported in literature, with large fenestrations resulting in regurgitation by Clothrap WH and Warren AT.

**Lambl’s Excrescences**

In 1856, Lambl first described small filiform processes on the ventricular surfaces of normal and abnormal aortic valves. In subsequent study of normal valves, a 70% to 90% prevalence of excrescences was reported, predominantly on the mitral (70-85%) and aortic valves (62-90%), but also on the right sided valves. They are seen in subjects 1-60 years old, regardless of gender. Valve excrescences consist of a core of connective tissue covered by endothelium.
Aetiology is uncertain. The constant movement of the cusps may lead to tearing of subendothelial collagen and elastic fibres, which subsequently endothelialize. This may explain their predominance on the left sided valves owing to large pressure differences. But the presence of excrescences on normal valves in children and adults of different age groups suggest that they may not result from an injury or age related process.

Valve excrescences have been reported with a frequency of up to 22% in patients with suspected cardioembolism referred for TEE. This finding has led to the suspicion that they are a substrate for embolism. Carlos A Roldan et al conducted a study on this topic and their findings are as follows: excrescences are common on the left heart valves, have a similar prevalence in normal subjects and in patients with or without suspected cardiac embolism regardless of age and gender and do not appear to be associated with embolic events.

In the present study, excrescences were seen on mitral, tricuspid and aortic valves; they were not seen in pulmonary valves. Excrescences were seen only in adults (27-88 years) and there was a male predominance (20 males and one female).

CONCLUSION
Knowledge about the dimensions and morphology of aortic valve is essential for diagnosis, prognosis and planning surgical procedures. It will also be helpful in manufacturing prosthetic valves.

Because great majority of patients with QAV and asymmetric cusps can develop complications, careful clinical evaluation and follow up is necessary. As there is a heritable component in BAV, all first-degree relatives of a patient with BAV should be screened by echocardiography. Patients with BAV should have healthy lipid profiles and must stop smoking to prevent early onset of stenosis.

Patients with large fenestrations on the cusps can develop rupture of cusps or AR leading to acute LVF. Surgical repair can be done to prevent such complications.

REFERENCES