The most common congenital abnormality of the aortic valve is bicuspid aortic valve, with an estimated incidence of about 2% of the general population [1]. Of note, anatomically isolated congenital valvular aortic insufficiency commonly occurs with bicuspid aortic valves, uncommonly with congenitally malformed trileaflet aortic valves (cuspal inequality) or unicommissural valves, and rarely with isolated quadricuspid aortic valves (QAVs). We report one patient who had QAV with aortic insufficiency, who presented with exertional dyspnea, and to and fro heart murmurs for 1 year.

CASE PRESENTATION

A 65-year-old male patient, who had a past history of chronic obstructive pulmonary disease, presented with dyspnea on exertion for a year. On physical examination, a 3/6, to and fro diastolic murmur over the left sternal border and expiratory wheezing were audible. Chest roentgenography showed cardiomegaly. Trans-thoracic echocardiography disclosed severe aortic insufficiency without valve sclerosis. Meanwhile, left ventricular end diastolic and systolic dimensions were determined to be 63.3 mm and 39.2 mm. Aortic root angiography demonstrated four aortic valve cusps with nearly equal size and aortic insufficiency jet (Figure 1). No other abnormality was documented at this time.

Thereafter, he underwent elective aortic valve replacement with 21 mm St. Jude Medical mechanical valve (St. Jude Medical, Inc., St. Paul, MN, USA) without incident. During surgical exploration, thin and morphologically symmetrical cusps with four commissures were confirmed (type A, according to the classification of Hurwitz and Roberts) (Figure 2) [2]. On histologic examination, the resected cusps showed fibrotic thickening with slight calcification and no sign of previous inflammatory disease. The patient underwent follow-up 8 months later and the postoperative echocardiography showed improved left ventricular end diastolic and systolic dimensions of 42.0 mm and 25.5 mm.
DISCUSSION

The first case report of QAV, carrying the predicted incidence of 0.008%, was published by Balington in 1862 [3,4]. Moreover, it has been estimated to have an incidence of about 1% among patients with pure aortic insufficiency [5]. The incidence of QAV was about 1.3% (2/140) among the surgically treated patients with pure aortic insufficiency at our institution, which is similar to the reported data. At present, it is estimated that more than 200 cases of QAV have been reported [6].

In reviewing the available data on QAV, most of them were discovered after the age of 40 (81%, 127/157 patients). This means that QAV could permit larger numbers of patients with this disorder to reach post-adolescence or even adulthood.

For the pathogenesis of QAV, several mechanisms have been proposed, such as excavation of one of the valve cushions or septation of a normal valve cushion from inflammatory process [7,8]. Nevertheless, the definite pathophysiology of QAV causing aortic insufficiency remains controversial. The anatomical anomaly of the cusps might induce unequal shear forces, which give rise to malcoaptation of the leaflets and result in aortic insufficiency [9]. On the other hand, the frequency of valvular dysfunction does not seem to be related to its unique morphology.

Most cases have been found incidentally, and presently QAV is increasingly being identified with the advent of echocardiography, transthoracic or transesophageal [10]. Therefore, a comprehensive investigation for other cardiac abnormalities is required since QAVs have also been discovered in patients with other cardiac abnormalities such as malposition ostium of coronary arteries, hypertrophic cardiomyopathy, or congenital anomalies of the other cardiac valves [11]. Fortunately, no obvious abnormalities were encountered in our patient and so the valve replacement was conducted simply. Furthermore, patients with QAV should be followed closely and surgical intervention should be done as early as possible for those with symptoms secondary to valvular insufficiency.

Symptomatic valvular insufficiency and correction of concomitant anomalies are the surgical indications for patients with QAV. Concerning the QAV, valve replacement remains the mainstay currently; nonetheless, surgical repair is technically achievable in a few selected patients [12]. However, no definite boundary exists between valve replacement and valvoplasty and more studies and long-term follow-up are needed in the future. Because of its vulnerability, proper antibiotic prophylaxis for possible endocarditis is advised before invasive procedures that may cause transient bacteremia [8].

In conclusion, QAV (which is a rare anatomic entity) needs to be taken into consideration, especially in patients without obvious documented etiology. Moreover, it must be considered as a malformation capable of leading to severe valve failure in later life.
and a close follow-up with echocardiography is warranted.

REFERENCES

罕見的四心瓣尖主動脈瓣解剖構造異常
合併主動脈瓣逆流 — 病例報告

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四心瓣尖的主動脈瓣是一種罕見的先天性心臟解剖構造缺陷，常在成人時期造成有症
狀的主動脈瓣逆流，進而需要手術置換；我們報告一位有四心瓣尖主動脈瓣合併主動
脈瓣逆流的病患，之後成功地接受主動脈瓣置換手術。

關鍵詞：主動脈瓣逆流，四心瓣尖主動脈瓣

(高雄醫誌 2007;23:422－5)