Normal coronary artery anatomy is characterised by two ostia centrally placed in the right and left sinus of Valsalva. The main left coronary artery (LCA) originates from the left ostium, branching into the left anterior descending artery and circumflex artery, which courses around the left atrioventricular groove; the right coronary artery (RCA) arises from the right ostium, providing an infundibular branch to the anterior side of the heart, and then courses backward in the atrioventricular groove. The three main coronary arteries branch superiorly to the atria and inferi orly to the ventricles; they end in broom-like arborisations, which penetrate the myocardium.1 Because of ventricular contraction myocardial perfusion of the left ventricle occurs mainly in diastole, while the myocardium of the right ventricle is perfused during both heart cycles.2–4

Clinical suspicion that a patient’s problems may be the result of coronary anomalies remains an important challenge in diagnosis, especially in children. Haemodynamically significant congenital anomalies of coronary arteries occur as isolated or primary forms and as secondary forms in association with congenital heart disease (CHD) (pulmonary atresia with intact interventricular septum or hypoplastic left heart syndrome with aortic atresia and severe mitral stenosis) (table 1). In this article only isolated/primary forms will be discussed.

INCIDENCE

Overall, anomalies of the coronary arteries are rather rare and the incidence of primary congenital coronary anomalies varies from 0.3% in a necropsy series reported by Alexander and Griffith to 1.6% of patients undergoing cardiac catheterisation in a series of more than 38 000 patients.5 Although these anomalies are rare, they may be seen with haemodynamic or myocardial perfusion abnormalities or high risk anatomy for accelerated atherosclerosis; they may result in symptoms varying from dyspnoea to sudden death. The milder forms escape detection both during life and at postmortem examination.

CLASSIFICATION

The classification of primary forms of coronary artery anomalies varies between ectopic origin of the coronary arteries from the aortic sinus, stenosis of the coronary arteries, absence of a coronary artery, to anomalous origin of the left or right coronary artery from the pulmonary artery or congenital coronary artery fistula.

SYMPTOMS

Although the spectrum of symptoms and associated syndromes is broad, the coronary artery anomalies that give rise to symptoms are limited to those that cause significant alteration in myocardial perfusion or result in pronounced left-to-right shunting (steal). The coronary anomalies most likely to cause myocardial infarction, ischaemia, or ventricular tachycardia are anomalous origin of the left coronary artery from the pulmonary artery (ALCAPA), large coronary arteriovenous fistulas, and those anomalies associated with a coronary artery coursing between the great vessels. The other coronary artery anomalies are rarely associated with symptoms or sudden death.

ANOMALOUS ORIGIN OF THE LEFT CORONARY ARTERY

Anomalous origin of the left main coronary artery (LMCA) (fig 1), the left anterior descending or circumflex branch from the proximal pulmonary artery or more distally to the proximal right pulmonary artery is a rare congenital anomaly occurring in 0.26% of patients with congenital heart disease undergoing cardiac catheterisation.4 In 1933 Bland, White and Garland described the clinical syndrome associated with this anomaly based on their experience with a 3 month old infant who died from it.7

When the RCA originates from the pulmonary trunk (ARCAPA), an anomaly rarer than ALCAPA, there is no ischaemia because the right ventricle is under low pressure; without
ischaemia there is no stimulus to form collateral anastomoses with branches of the LCA and thus there is no great tendency to establish fistulous flow.

Both coronary arteries arising from the pulmonary trunk constitute an extremely rare situation. In such cases symptoms appear within a few days of birth, and death follows within two weeks; it is compatible with life only if associated with pulmonary hypertension.

Since Edwards' fundamental contribution, it has been known that dramatic changes occur in the coronary circulation in all children with ALCAPA within the first weeks of life. While the fetus remains in utero, the heart develops quite normally. After birth, as long as the pulmonary arterial pressure remains at or near systemic levels, the left ventricular myocardium supplied by the anomalous artery remains well perfused; the avidity of myocardium for oxygen is such that normal perfusion with blood of mixed venous oxygen content does not present a problem.

As pressure in the pulmonary trunk falls postnatally, perfusion of the left ventricle begins to suffer because the period during which pressure in the coronary artery exceeds intramural left ventricular pressure becomes shorter. The consequence of the circulatory handicap is a decrease of left ventricular end diastolic pressure with consecutive pulmonary vasoconstriction; slowing of the postnatal reduction of pulmonary artery pressure and migration of the perfusion problem will result. At the same time, the ischaemic myocardium is being perfused increasingly by a developing set of collateral vessels from the RCA which arise normally.

Thus this malformation postnatally presents a picture which may vary enormously from case to case and from time to time in a given patient (table 2).

The small infant usually presents with signs of congestive heart failure and a particular type of anginal attack, while the older child and adolescent present with an unexplained heart murmur caused by mitral insufficiency, mild cardiomegaly, or an abnormal ECG. Myocardial ischaemia may present first in adolescence or young adult life when, under stress or maximal motivated exertion, either anginal pain or arrhythmia may occur. The latter may result in sudden unexplained death or near death. While the ECG is not likely to be normal, it may not show the classical findings in ischaemia or infarction, so that there are occasions with this disease when the ECG is not diagnostic. The older child who has survived without symptoms may very well have a normal ECG.

Table 1 Haemodynamically significant congenital anomalies of the coronary arteries

<table>
<thead>
<tr>
<th>Isolated/primary—without CHD</th>
<th>Secondary—with CHD</th>
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<tr>
<td>▶ Anomalous origin of accessory coronary arteries from the pulmonary artery (ALCAPA, ARCAPA)</td>
<td>▶ PA + IVS</td>
</tr>
<tr>
<td>▶ Ectopic origin of the coronary arteries from aortic sinus</td>
<td>▶ AA + MS</td>
</tr>
<tr>
<td>▶ Absence of a coronary artery</td>
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<tr>
<td>▶ Congenital coronary artery fistula</td>
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AA, aortic atresia; ALCAPA, anomalous origin of the left coronary artery from the pulmonary artery; ARCAPA, anomalous origin of the right coronary artery from the pulmonary artery; CHD, congenital heart disease; IVS, interventricular septum; MS, mitral stenosis; PA, pulmonary atresia.

Table 2 Symptomatology of ALCAPA

- Perfectly distributed collateral arteries: minor ischaemic damage
- Collaterals poorly distributed: some parts well perfused/some ischaemic
- Large interconnections: steal

Three groups of patients can be identified:

- those with perfectly distributed collateral arteries before the pulmonary artery pressure had fallen with minor ischaemic damage of the left ventricular myocardium, which is well perfused, and normal in form and function; this type is often diagnosed in adolescents or adulthood
- those with poorly distributed collateral connections—some parts of the left ventricle are well perfused while others become ischaemic
- those with collateral maldistribution, where there are large interconnections proximal to the branches supplying contracting myocardium which act as a left-to-right arterial shunt, requiring extra work for the left ventricle while diverting blood supply from the myocardium (steal from the myocardium to the pulmonary artery).
the coronary arteries. An aortic root injection may be sufficient to show the early filling of the RCA and the delayed passage via collateral vessels into the LCA and finally the pulmonary trunk.

The finding of coronary artery filling from the pulmonary trunk should rule out ligation of the LMCA at its origin as a form of treatment for the patient.

The diagnosis of ALCAPA in an infant, usually seriously ill, is an indication for urgent surgery; diagnosis in older patients should also be an indication for surgery. Excellent surgical results have been reported following re-establishment of a dual coronary artery system, with direct reimplantation of the coronary artery into the aorta, left subclavian artery to LCA anastomosis, or transpulmonary baffling or Takeuchi procedure, as it restores oxygenated blood flow to the left ventricular myocardium immediately and more completely. Although clinical improvement with simple ligation of the LMCA may rise to the right of the RCA and pass posteriorly to the aortic root.

The origin of the LMCA from the right sinus of Valsalva can be classified into four types (fig 4):

- the LMCA passes between the aorta and pulmonary trunk, posteriorly and adjacent to the pulmonary trunk
- the LMCA passes anteriorly over the right ventricular outflow tract
- the LMCA may course along the crista supraventricularis intramyocardially or subendocardially, surfacing in the proximal interventricular sulcus
- the LMCA may rise to the right of the RCA and pass posteriorly to the aortic root.

Cheitlin and his colleagues have pointed out the pathologic significance of a single coronary artery or of both coronary arteries from the right sinus of Valsalva when the
branch or artery that supplies the left coronary distribution courses leftward between the aorta and pulmonary trunk so that it can be compressed (fig 5). They showed a striking incidence of sudden unexplained death in adolescents and young adults, and a strong relation of death to heavy exertion. For this reason, any child or young person with angina pectoris, myocardial infarction, or cardiac syncope should have this anomaly ruled out by coronary angiography.

In contrast, when a single coronary artery arises from the left sinus of Valsalva or both coronaries arise from separate ostia in that sinus with the RCA supply coursing between aorta and right ventricular outflow tract, compression can occur but sudden death usually does not, probably because of the low perfusion pressure required for the right ventricle.

An interesting finding was published by Bellhouse and colleagues in 1968: they reported that the coronary blood flow is significantly attenuated if the orifice of the coronary arteries lies above the sinus of Valsalva beyond the sinotubular junction. This finding should be taken into consideration in patients with transposition of the great arteries after the arterial switch operation with reimplantation of the coronary arteries into the ascending aorta.

Indications for surgery remain debatable; however, reversible ischaemia in the anomalously connecting artery is an indication. Coronary artery bypass grafting, using the internal thoracic artery graft to the anterior descending artery and a saphenous vein graft to the circumflex artery, is feasible in patients with anomalous connection of the LMCA to the right sinus of Valsalva. Alternatively, the intramural part of the coronary artery can be opened longitudinally from within the aorta, with reconstruction of the most proximal part of the coronary artery, so that the ostium emerges from the left sinus of Valsalva in a normal and unobstructed location. The absence of the LMCA was noted as the most

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**Figure 4** Origin of the left main coronary artery (LMCA) from the right sinus of Valsalva. Top left: the LMCA passes between the aorta and pulmonary trunk, posteriorly and adjacent to the pulmonary trunk. Top right: the LMCA passes anteriorly over the right ventricular outflow tract. Bottom left: the LMCA may course along the crista supraventricularis intramycardially or subendocardially, surfacing in the proximal interventricular sulcus. Bottom right: the LMCA may rise to the right of the right coronary artery and pass posteriorly to the aortic root.

**Figure 5** Coronary artery compression.
Congenital arteriovenous/arterio-arterial fistulae

Congenital arterio-venous or arterio-arterial fistulae (CAF) of the coronary arteries are direct connections between a branch of a coronary artery and the lumen of the four cardiac chambers, the coronary sinus, the superior vena cava, the pulmonary artery, or the pulmonary veins close to the heart without the intervening capillary system. The source is most often the RCA with the LCA being much less involved. Embryologically, these fistulae seem to represent persistent junctions of primordial epicardial vessels with intramyocardial sinusoidal circulation. Most patients present late in life, occasionally in childhood and rarely in infancy.

Symptoms of the disease may be a continuous systolic-diastolic murmur, congestive heart failure, pulmonary hypertension, bacterial endocarditis in 5% of cases, premature arteriosclerotic changes within the fistulae, and thromboembolic events. Angina is uncommon and myocardial infarction rare; it is postulated that these ischaemic symptoms are caused by coronary steal. CAF that have not been detected or closed in childhood have been reported to become symptomatic in adulthood because of chronic volume load and ischaemia. Thus, it has previously been recommended that these fistulae be closed in childhood, either surgically or, as suggested more recently, by transcatheter coil occlusion. Incidental detection of CAF in asymptomatic patients has been noted on coronary angiography, but their clinical significance is unclear.

With the advent of high resolution two dimensional and colour Doppler echocardiography during the past decade, the incidental diagnosis of clinically silent CAF is increasing. Colour flow mapping has proved to be a highly sensitive method for identifying small CAF (fig 6). Cardiac catheterisation and selective coronary angiography are necessary for definitive diagnosis and planning of either surgical repair or occlusion by interventional catheterisation (fig 7).

Currently, there is a paucity of information regarding the clinical implications of incidentally identified, clinically silent CAF. Some authors have recommended elective closure of CAF regardless of symptoms, while others have advocated conservative management. The current general opinion does not support the previous recommendations to close CAF even in asymptomatic patients; a compelling argument in favour of a conservative approach to these patients is the unexpectedly high incidence of spontaneous closure of CAF.

Several clinically relevant questions remain unanswered. The lifelong risk of complications such as infective endocarditis, aneurysm formation, dissection, rupture, accelerated atherosclerosis, and thromboembolism is not known. Nevertheless, because any intervention to close such fistulae is associated with a risk of morbidity, and because the benefit of such intervention is questionable, a conservative approach to small asymptomatic CAF is recommended.

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Congenital anomalies of the coronary arteries

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