Case reports

Recurrent ventricular tachycardia associated with anomalous left coronary artery from the pulmonary artery in a child managed by revascularisation and map-guided endocardial resection

JANET M McCOMB, RICHARD VINCENT,* COLIN J HILTON

From the Regional Cardiothoracic Centre, Freeman Hospital, Newcastle upon Tyne and *Royal Sussex County Hospital, Brighton

SUMMARY A 13 year old girl with recurrent ventricular tachycardia that occurred in association with anomalous origin of the left coronary artery from the pulmonary artery is described. Such an arrhythmia has not been described before. Ligation of the anomalous left coronary artery led to more frequent episodes of ventricular tachycardia with evidence of myocardial ischaemia. She was successfully managed by revascularisation (to establish a two coronary system) and map-guided left ventricular aneurysmectomy.

Anomalous origin of the left coronary artery arising from the main pulmonary artery is a rare congenital anomaly that is often fatal in infancy. Myocardial ischaemia is common and infarction may occur, as may mitral regurgitation. The commonest mode of presentation is cardiac failure in infancy, but sudden death may occur both in children and adults with or without previous symptoms. In a review of 147 patients, sudden death was reported in 11, occurring after exertion in six. Eight had been symptom free. Sudden death in these patients is thought to be caused by ventricular arrhythmias, although such arrhythmias have rarely been recorded.

Management is controversial, with both medical and surgical treatment giving poor results, although a comparison was difficult because of bias in the selection of patients. There have been various surgical approaches, although evidence suggests that revascularisation, with establishment of a two coronary system, is the best option.

We describe a 13 year old girl who is unusual in several respects. She has survived to adolescence, despite a mortality for this condition in infancy of 80-90%. She presented with recurrent ventricular arrhythmias, which increased in frequency after ligation of the anomalous left coronary artery. Further recurrence of arrhythmias was prevented by surgical revascularisation with map-guided ventricular aneurysmectomy.

Case report

This 13 year old girl presented initially at the age of three months when she became cyanosed and dyspnoeic. She was found to be in heart failure with tachycardia, cardiomegaly, and hepatomegaly. She improved gradually after treatment with digoxin, which was subsequently stopped.

She remained well until the age of 11 when during
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gymnastics she had an episode of ventricular tachycardia with syncope, chest pain, and dyspnoea (fig 1). After she was admitted to her local hospital an apical systolic murmur was heard. The electrocardiogram showed Q waves in leads I and aVL with loss of R wave amplitude and non specific ST-T wave changes in the lateral chest leads (fig 2). The chest x ray showed moderate cardiac enlargement and the echocardiogram showed "slight dilatation of the left ventricle with some hypokinesia." Atenolol 25 mg daily was started and later was increased to 50 mg twice a day. Five months later, after a sudden fright she had a further episode of tachycardia, which terminated spontaneously after 45 minutes. After this, she was able to exercise on the treadmill according to the standard Bruce protocol for 14

Fig 1 Electrocardiogram recorded during first episode of ventricular tachycardia.

Fig 2 Electrocardiogram recorded during sinus rhythm.
minutes, the heart rate rising from 67 to 166 beats per minute. Asymptomatic ST depression developed in the inferior leads and resolved during recovery. Ventricular bigeminy was seen at the termination of exercise. One week later, she complained of mild discomfort in the left arm, legs, and chest, with muscle pains and headache. She went swimming, and after two lengths tachycardia and clenching chest pain developed, which radiated to the back and was eased by glyceryl trinitrate. Runs of ventricular tachycardia were subsequently seen.

She was then referred for invasive studies. Cardiac catheterization showed an anomalous left coronary artery arising from the main pulmonary artery. The diagnosis was confirmed at operation, when the left coronary orifice was oversewn from within the pulmonary artery. The heart was enlarged and there was a large anteroapical fibrous left ventricular aneurysm. On day 7 an episode of sustained ventricular tachycardia developed and she was treated with flecainide 100 mg twice a day. Three weeks after this she had an episode of palpitation lasting two hours caused by ventricular tachycardia. This was treated by cardioversion. Treatment was changed to mexiletine 100 mg twice a day. Three weeks later she felt "a run of thumps" while cycling. She dismounted, went indoors, and fainted. Her mother described her as deep purple, with a few gasping respirations. She lay still, had no pulse, and no heart beat. Cardiopulmonary resuscitation was started and she recovered within 30 seconds.

She was then transferred to Freeman Hospital. By this time she complained of interscapular pain on exertion and was afraid to exercise. Dipyridamole-thallium scintigraphy confirmed a reversible perfusion defect in the anterior wall, and a coronary angiogram showed a large tortuous right coronary artery. The left coronary artery opacified quickly after injection of the right coronary artery, but its filling was less pronounced than it had been before ligation. No ventricular arrhythmias were induced by programmed ventricular stimulation. At operation, epicardial mapping of both ventricles was performed during normothermic bypass. This identified an area of abnormal electrical activity associated with the apical aneurysm. The left ventricle was then opened through the apical scar and endocardial mapping was performed during sinus rhythm. This confirmed an area of abnormal electrical activity in the aneurysm. Programmed ventricular stimulation again failed to induce ventricular arrhythmias. The endocardium from which fractionated electrograms were recorded was resected, as was the aneurysm. The left internal mammary artery was grafted to the left anterior descending coronary artery during cold cardioplegia.

Postoperatively she made a good recovery. Subsequently, she has done well, growing 11 cm within three months. She is taking no medication, is exercising normally, has no angina, and there has been no recurrence of her arrhythmia.

Discussion

Malignant ventricular arrhythmias are a well recognized sequel of ischaemic heart disease caused by coronary atheroma, but have been less well documented in other forms of myocardial ischaemia. Sudden death has, however, been associated with an anomalous left coronary artery arising from the pulmonary artery, and there is evidence that this is more common in adults. Although it has been suggested that arrhythmias are the cause of sudden death in these patients, such arrhythmias have rarely been described in association with this anomaly.

We report an unusual presentation of anomalous left coronary artery, with recurrent sustained ventricular tachycardia and syncope occurring for the first time in adolescence. Heart failure in infancy had resolved with medical treatment, and the patient had been symptom free for 11 years, although there was evidence of silent myocardial ischaemia. Spontaneous resolution of symptoms has been described but is unusual. Her arrhythmias increased in frequency after ligation of the anomalous artery, and ischaemia was again demonstrated. All her symptoms resolved after endocardial resection and revascularisation with the left internal mammary artery.

There has been controversy about the most appropriate treatment, with some advocating medical treatment and some surgical treatment. Various surgical procedures have been recommended, ranging from ligation of the anomalous artery to revascularisation procedures, including direct re-implantation, saphenous vein grafting, and arterial flap repairs. Despite this, surgical results have been described as disappointing. Simple ligation of the anomalous coronary artery is associated with a high mortality, which has been attributed to ischaemia in the absence of collateral circulation. Ligation leaves a single coronary system, which may lead to subsequent ischaemia. Angina developing some time after ligation has been reported. Asymptomatic ischaemia has also been shown by exercise testing in approximately half of the few patients so studied. Ligation may also be associated with a higher risk of late sudden death than other surgical procedures. Revascularisation, with the establishment of a two coronary system, is therefore thought to be the best treatment. Attempts at establishing a two coronary system have had varying results. Early attempts used saphenous vein grafting to the anomalous left
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coronary artery after ligation. Over 30 patients with saphenous vein grafts have been reported. Grafts were shown to be patent postoperatively in 11 patients, and were known to be occluded in five (31%). Various other methods of revascularisation have been described, but none was appropriate in our patient because of the previous ligation. The left internal mammary artery was therefore used to graft the anomalous coronary artery. This technique, not previously described in this condition, satisfactorily established a two coronary system, and may remain patent for longer than a saphenous vein graft.

Map-guided left ventricular endocardial resection was performed in addition to bypass grafting in our patient, as a further antiarrhythmic measure. Others have performed aneurysctomy or left ventricular resection in a few patients, presumably because of heart failure, but a map-guided procedure has not been described before.

This report describes ventricular tachycardia in association with an anomalous left coronary artery, and emphasises the importance of appropriate management of ischaemia in the treatment of this condition.

References


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doi: 10.1136/hrt.62.5.396