CONGENITAL CORONARY ARTERY–RIGHT VENTRICULAR FISTULA CORRECTED SURGICALLY*

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Anomalous communication of one or both coronary arteries and a cardiac chamber was described for the first time by Cayla in 1885. Recently, the introduction of selective angiocardiography and the possibility of satisfactory repair by relatively simple surgical procedures have furthered the interest in the syndrome. Most of the patients diagnosed during life and operated upon were children (Gasul et al., 1960; Abbott, Rivarola, and Logue, 1961; Cooley and Ellis, 1962; Soulié et al., 1963; Michaud et al., 1963; Effler and Welti, 1963).

The patient presented here is unusual in that he was a middle-aged man with a congenital fistula between the right coronary artery and the right ventricle, which was successfully treated by operation.

Case Report

A 33-year-old white accountant was admitted to Chicago Wesley Memorial Hospital in May 1961, for evaluation of a heart murmur and cardiomegaly, which had been present since birth. His past history revealed an episode of subacute bacterial endocarditis in 1946, successfully treated with penicillin in another hospital. In 1955 and again in 1956, he was admitted to the hospital for "kidney infection", and in 1957 he underwent right inguinal hernia repair. The patient had been entirely asymptomatic throughout his life and had never limited his physical activity. However, he was not accepted for military service, and life insurance had been denied to him because of his cardiac status.

On physical examination, he appeared well developed, obese, not in acute or chronic distress. The respiratory rate was 20 per minute, pulse rate 76 per minute, and blood pressure 130/70 mm. Hg. There was no precordial asymmetry. The point of maximal impulse was strong and diffuse and was located at the sixth intercostal space at the anterior axillary line. The cardiac dullness extended 3-5 cm. to the right of the sternum. There was a significant left parasternal heave and a systolic thrill was palpable at the third left intercostal space at the sternal margin. In the same area, a loud (grade 4 out of 6) systolic and an early-to-mid medium intensity (grade 3 out of 6) diastolic murmur were present, which radiated widely over the anterior præcordium. The second sound was normally split. The peripheral pulses were normal.

The chest radiograph showed increased pulmonary vascularity. The transverse cardiac diameter was 42 per cent above the normal and the enlargement appeared to be biventricular. There was marked increase in the diameter of the main pulmonary artery segment. The electrocardiogram showed incomplete right bundle-branch block pattern, and there was left axis deviation (−15°) and increased voltage in the precordial leads (RV6+SV1=46 mm.) suggesting left ventricular hypertrophy (Fig. 1). The phonocardiogram confirmed the presence of the systolic and diastolic murmurs. Right heart catheterization disclosed a significant increase in oxygen content at the ventricular level. The left-to-right shunt was 3.9 l./min./m.² with a systemic output of 2.3 l./min./m.². The right ventricular systolic pressure was moderately raised (43/2 mm. Hg) and a gradient of 23 mm. Hg was present across the pulmonary valve: this was interpreted as due to the increased flow. Total pulmonary resistance was 441 dynes sec./cm. −5. Retrograde aortography from the right brachial artery with injection into the left ventricle showed no ventricular septal defect but instead demonstrated a very dilated right coronary artery and ascending aorta (Fig. 2).

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781
Operation was performed on May 10, 1962. The pulmonary artery and the heart, as seen through the pericardial sac, appeared exceedingly tense. On opening the pericardium, the right coronary artery was noted to be extremely dilated, measuring approximately 4 cm. in diameter. Under total cardiopulmonary bypass, a right ventriculotomy was performed and it was noted that the dilated coronary artery was draining into the apical region of the right ventricle. However, due to the many recesses present in this area, the exact location of the opening could not be detected from the ventricular side. Therefore, after cross-clamping the aorta, the coronary artery was opened longitudinally and the opening of the fistula easily visualized. A probe was inserted through the opening. Closure of the fistula was accomplished with two layers of 5-0 silk, and the continuity of the coronary artery was reinstituted. The aorta was cross-clamped for a total of 40 minutes.

The patient made an uneventful recovery. The electrocardiogram showed the usual post-operative subepicardial injury pattern. The patient returned for observation eight months after operation. By this time, he had resumed full activity and was symptom free. No murmurs were detected on physical examination. The post-operative electrocardiographic changes had disappeared, and the voltage in the precordial leads was now normal. The chest radiography showed a decrease in the transverse cardiac diameter to 22 per cent above the normal.

Discussion

According to Edwards (1958) the anomalies of the coronary arterial system may be divided into three groups: (1) minor anomalies of origin or distribution of no importance under usual circumstances; (2) anomalies secondary to other cardiac malformations; and (3) primary anomalies of origin or of communication that are of major significance. In this group, maldevelopment may produce two types of abnormal arteriovenous communications. In the first, a coronary artery originates from the pulmonary trunk with reversal of the blood flow in this vessel which is fed by anastomosis from the other coronary artery (Edwards, 1958). In the second variety, an arrest or
faulty differentiation of the embryonal intramyocardial sinusoids is responsible for the communication between a coronary artery and a right or left-sided cardiac structure (Edwards, Gladding, and Weir, 1958). In this report, we are concerned only with the second variety of Edwards' third group of coronary arterial abnormalities, since the pathology in our patient consisted of an anomalous communication between the right coronary artery and the right ventricle.

The primary functional disturbance of a coronary artery fistula is the "run-off" of blood from the coronary arterial system, with the portion of myocardium supplied by the vessels beyond the communication receiving less blood than normal. In fact most of the blood is diverted away from the myocardial capillaries, which represent a zone of high resistance, to the low-resistance anomalous channel. However, one can assume that an efficient collateral circulation supplying the critical area develops early in life since myocardial ischaemia as a complication of the syndrome has been reported in only a few instances (Knoblich and Rawson, 1956; Valdivia, Rowe, and Angevine, 1957; Edwards et al., 1958) and never in infants. Our patient never complained of angina. The transitory electrocardiographic changes after operation are a common finding in operations involving the heart and are probably due to pericardial injury and not secondary to this anomaly. The majority of the cases with coronary artery–right ventricular fistula reported by others have had continuous murmurs. The diastolic component was reported to be louder in 4 out of 5 patients of Gasul et al. (1960). Separate systolic and diastolic murmurs have also been described. Since it is expected that flow through the fistula is greatest during diastole, the diastolic component of the murmur should be louder. However, in our case, the systolic component was louder. It appears that there was a superimposed ejection murmur which made the systolic murmur more prominent. This explanation is supported by the fact that there was a 23 mm. Hg systolic gradient at the pulmonary valve during cardiac catheterization. Since the fistula entered the apical region of the right ventricle, shunting was probably less during systole because of narrowing of the fistulous opening during contraction of the myocardium. Though the patient was asymptomatic, his right ventricular systolic pressure and total pulmonary resistance were moderately raised due to the large left-to-right shunt. In consequence of the overload of all cardiac chambers, the heart was very enlarged, including the pulmonary trunk. Dilatation of the right coronary artery and of the ascending aorta was well demonstrated by selective angiography. This latter feature has been emphasized by Gasul et al. (1960) as one of the leading characteristics of the syndrome, short of the demonstration of the dilated coronary artery draining into a cardiac chamber.

Patients with this condition seem to tolerate the increased cardiac load (which in some cases may be quite large) extremely well (Gasul et al., 1960; Neufeld et al., 1961; Carmichael and Davidson, 1961). However, recently Steinberg, Baldwin, and Dotter (1958) described 5 adult patients who died in congestive heart failure produced by a similar anomalous communication. Congestive heart failure may be a prominent feature also in infancy (Sanger, Taylor, and Robicsek, 1959; Cooley and Ellis, 1962; Braudo et al., 1962) and be the cause of death. Other complications include pulmonary hypertension (Davison, McBracken, and McIlvain, 1955; Neill and Mounsey, 1958; Bosher et al., 1959) and bacterial endoarteritis (present case; Trevor, 1911; Jacobi and Heinrich, 1933; Vela, Velasques, and Fuenmoyor, 1951; Sanger et al., 1959). Therefore, the syndrome does not appear to be as benign as the early investigators thought.

The current view regarding therapy holds that the presence of either clinical manifestations or significant laboratory findings calls for surgical repair of the lesion. This has been accomplished in 41 patients to date (Sondergaard, 1955; Davis et al., 1956; Johnson, personal communication to Davis et al., 1956; Mozen, 1956; Morrow, personal communication to Edwards et al., 1958; Neill and Mounsey, 1958; Bosher et al., 1959; Grob and Kolb, 1959; Kittle, 1959; Sanger et al., 1959; Swan et al., 1959; Diehl, 1960; Gasul, et al., 1959; Zuhdi et al., 1960; Carmichael and Davidson, 1961; Dubost, Chevrier, and Metianu, 1961; Engle et al., 1961; McIntosh et al., 1961; Neufeld et al., 1961; Barcia et al., 1962; Braudo et al., 1962; Cooley and Ellis, 1962; Papaioannou et al., 1962; Effler and Welti, 1963; Michaud et al., 1963; Soulié et al., 1963) with only 2 reported deaths (Bosher et al., 1959; Michaud et al., 1963): both occurred in small critically ill children, one of whom had an
associated patent ductus arteriosus. Three patients in whom interruption of the fistula was obtained by ligation of the coronary artery developed myocardial infarction (Swan et al., 1959; Cooley and Ellis, 1962; Michaud et al., 1963). In a fourth patient cardiac arrest occurred four hours after operation (Cooley and Ellis, 1962).

Therefore, closure of the fistula without interruption of the continuity of the parent vessel was advocated and has been successfully accomplished in this patient.

**Summary**

A case of right coronary artery–right ventricular fistula occurring in a middle-aged asymptomatic man is reported. Physiological study was prompted by the detection of an unexplained cardiomegaly and heart murmur. Correction of the lesion was accomplished by direct vision closure of the fistula with maintenance of the parent coronary artery patency. The patient made a complete recovery.

Recent publications on the subject and current therapeutic orientation are discussed.

The operation in this patient was performed by Dr. Arthur De Boer, staff surgeon of Chicago Wesley Memorial Hospital.

**References**


Johnson, J. Quoted by Davies et al. (1956).


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