Case reports


Antepartum diagnosis of vasa praevia—report of a case causing sudden foetal death

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Summary
A case of sudden foetal death due to vasa praevia is reported. The main problem is early antepartum diagnosis. Three methods are discussed.

1. Tests to recognize foetal blood as a routine in antepartum haemorrhage;
2. amnioscopy before amniotomy; in foetal irregularity or bradycardia; in antepartum haemorrhage;
3. tests for foetal bradycardia in early labour as a part of the routine abdominal examination.

Introduction
Vasa praevia is a rare obstetric complication which must be kept in mind as a rare cause of antepartum haemorrhage. Less than a hundred cases have been reported in the literature. It causes two symptoms—foetal bradycardia when a vessel is compressed, and a haemorrhage when the vessel is torn. Severe bradycardia will lead to Caesarean section for foetal distress of probably 'unknown' origin—the infant may be saved, the cause of the foetal distress only being apparent on inspection of the placenta after the operation.

Haemorrhage is dangerous to the foetus because the blood-loss is of foetal origin. What would be considered an insignificant haemorrhage if the blood-loss was maternal in origin would be lethal to the infant when foetal blood was lost—as illustrated by this case, where the foetal heart failed with the very first bout of bleeding. One difficulty in the management of antepartum haemorrhage is to exclude the possibility of the blood-loss being of foetal origin.
Case report

A 39-year-old woman, Gravida IV Para III. Three previous pregnancies uneventful, the last baby being born 12 years previously.

Preliminary investigations

WR and GCFT, negative. Blood group, O Rh. negative. Hb, 75%. BP, 120/60. Nothing abnormal detected in cardiovascular and respiratory systems. Pregnancy was uneventful until she was admitted in labour at the thirty-seventh week of gestation. Her general condition was satisfactory. Blood pressure 110/70. On examination the height of the fundus was near term, presentation vertex-engaged. Internal examination revealed the cervix was taken up, os was 2 fingers dilated, membranes were intact. Vertex was engaged.

Three hours after admission, whilst in labour she had a brisk bleed of about 100 ml. The foetal heart was found to be absent almost simultaneously. Bleeding ceased after a few min. The foetal parts were easily palpable. There were no tender areas over her uterus. The consistency of the uterus felt normal, which clinically ruled out any likelihood of accidental haemorrhage. Vaginal examination at this stage showed the os was 2 fingers dilated, vertex engaged, and membranes ruptured. As the foetal heart had failed, labour was allowed to progress to delivery. Fifteen hours after admission the third stage was completed by Brandt-Andrew manoeuvre. Inspection of the placenta revealed a low velamentous insertion of the cord with a torn major vessel (Fig. 1).

The cause of the haemorrhage and the sudden failure of the foetal heart were found to be due to rupture of the membranes and a torn major vessel. The presenting part had compressed the vessel to prevent further bleeding.

The infant, male, weighed 8 lb 6 oz; 22 in long; head circumference 13 in. No abnormality externally evident. Unfortunately, the parents refused permission for autopsy.

Discussion

The foetal heart stopped almost simultaneously with the short, brisk bleed. The problem of vasa praevia needs consideration. Once labour has started, the cervix opens leaving the membranes unsupported and rupture of the vasa praevia is almost inevitable. Once antepartum haemorrhage occurs, there is little hope for the foetus, unless the possibility of the blood being of foetal origin is recognized. Some workers advise the Apt test (Apt and Downey, 1955) to distinguish foetal from maternal blood. An alternative is the Kleihauer test.

Naftolin and Mishell (1963) reported three cases of vasa praevia with the loss of one infant, but the diagnosis was made only after delivery. Curl and Johnson (1968) reported one case in which the pulsation of the vessels was detected at routine vaginal examination: a Caesarean section saved the infant. Paulino (1970) reviewed the literature over a 10-year period and reported six cases with one live infant. Only in one instance was the vasa praevia diagnosed ante partum. Barham (1968) reported a case in which the vasa praevia was detected by amniocentesis; he recommends the procedure in the following instances: (a) before amniotomy (whatever the indication); (b) when foetal irregularity occurs in labour; (c) when unexplained ante-partum haemorrhage occurs.

If amniocentesis were done routinely in the above situations, it might be possible to make a diagnosis of vasa praevia before symptoms of foetal anoxia were manifest.

Inducing foetal bradycardia at early labour as a part of the routine abdominal examination may help to focus attention on those foetuses which are liable to get into distress. The foetal heart rate is taken between uterine contractions and then repeatedly during contraction while pressing the head well into the pelvis. If the rate falls steeply or the slow rate persists after the contraction has ceased, an extra careful watch is kept over the foetus. A presentation of the cord has been diagnosed by this method and could help in the ante-partum diagnosis of vasa praevia (Amarasinghe, 1970).

References


Traumatic aortic incompetence following road traffic accident

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Summary
This case report describes the presentation and treatment of a case of aortic incompetence, resulting from a road traffic accident. The relevant literature is briefly reviewed.

Aortic incompetence due to trauma has been described following non-penetrating chest injuries, such as kicks from horses (Barie, 1881), falls from heights and crushing accidents (Kissane, Koons and Clark, 1948; Levine, Roberts and Morrow, 1962). Despite the frequency of road traffic accidents, there have been no recent reports of traumatic aortic valve damage.

Case history
The patient, a 44-year-old transport manager, was admitted to hospital in September, 1972. He gave a history of increasing breathlessness on exertion over the previous 15 months. For the 2 months before admission he had complained of nocturnal dyspnoea. The symptom of breathlessness as well as episodes of dizziness had dated from May, 1971, when he had been involved in a serious road accident. The nature of the accident was such that, on impact, he had been thrown through the back window of his car, sustaining injuries to the upper thoracic spine and occipital part of the skull. He was unconscious for approximately 10 min and, on admission to the casualty department, the principal abnormality was of tenderness, swelling and marked angulation of the thoracic spine. X-rays revealed a compression fracture of the 7th thoracic vertebra. No murmurs were noted and blood pressure was normal. Systolic and early diastolic murmurs were first noted 5 days after admission.

Following discharge the patient noted that he was breathless on climbing stairs and that he suffered quite marked light-headedness on more severe exercise.

In the past history, he had a myocardial infarction in 1968. During that admission only a soft systolic murmur was noted. There were no signs of aortic insufficiency, the heart was of normal size and the electrocardiogram showed normal left ventricular voltages. He was said to have had rheumatic fever at the age of 28, treated at home. Subsequent medical examinations in the Services were satisfactory.

On admission to hospital, he had signs of left ventricular failure and of gross aortic incompetence. Good progress was made on standard therapy but shortly after discharge he was re-admitted in acute pulmonary oedema and subsequently transferred for more specialized investigation.

Examination
The patient was pale, mildly dyspnoeic with marked Corrigan pulsations in the neck. Jugular venous pressure was elevated and a third sound was audible on auscultation. There was a loud aortic systolic murmur and long aortic diastolic murmur, both radiating to the neck and maximal over the upper left sternal border. Pulse was collapsing, sinus in rhythm, and blood pressure was 180/50. There were bilateral fine crepitations in the chest.

Investigations
Chest X-ray showed cardiomegaly and pulmonary venous congestion and the ECG showed marked left ventricular hypertrophy. At cardiac catheterization, the left ventricular end diastolic pressure was 15–45 mmHg and a supravalvular angiogram showed severe aortic incompetence. Coronary angiograms were normal except for minor narrowing of the left anterior descending artery. WR was negative.
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