A case of sirenomelia-symphus monopus

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Introduction
Sirenomelia (mermaid deformity) consists of varying degrees of lower limb fusion and was first described by Rocheus in 1542 who named it after mythical Greek sirens¹. The worldwide incidence is 0.8-1 case in 100,000 live births, the male to female ratio being 3:1². It is usually associated with multiple anomalies of the gastrointestinal, genitourinary, cardiovascular and musculoskeletal systems². Here we describe a rare case of sirenomelia symphus monopus.

Case report
A 750g early preterm baby was born by spontaneous vaginal delivery to a 25 year old gravida 3 para 1 mother who had one abortion. The marriage was non-consanguineous and the mother was from a low socioeconomic class. She attended the antenatal clinic regularly and received iron, folic acid and calcium supplementation. Prenatal ultrasound was suggestive of severe oligohydramnios. She had no history of pregnancy induced hypertension, pre-gestational or gestational diabetes. There was no history of smoking or tobacco chewing. There was a history of first trimester abortion but there was no family history of fetal anomalies. Apgar score was 0 at first and fifth minutes of life. Respiratory efforts after birth were poor. The baby could not be salvaged and died within 30 minutes of birth.

On examination, baby had flattened facies, microcephaly, high arched palate, fused eyelids and malformed ears. Baby exhibited only the right upper limb with deformed forearm and hand and only the limb bud of left upper limb. The left lower limb demonstrated deformity of the leg and foot mimicking a tail with absent right lower limb and pelvic girdle (Figure 1).

Anal opening and external genitalia were absent, and baby also had a single umbilical artery. Radiograph of the baby showed absent radius in the right upper limb and absent fibula in left lower limb (Figure 2).

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Discussion
Sirenomelia was originally divided into 3 categories depending on the number of feet present: symphus apus (no feet), symphus monopus (one foot) and symphus dipus (both feet). This case belongs to the symphus monopus category. A better categorisation is that of Stocker and Heifetz in which seven types are defined: I (all thigh and leg bones present), II (single fibula), III (absent fibulae), IV (partially fused femurs, fused fibulae), V (partially fused femurs, absent fibulae), VI (single femur, single tibia) and VII (single femur, absent tibia). This case belongs to category VI. The aetiology of this anomaly remains unclear. Whilst death is the usual outcome, as in our patient, survivals have been reported. A study by Orioli et al found that 50% cases of sirenomelia were associated with genital, large intestinal, and urinary defects and 10-15% with lower spinal column defects, single umbilical artery, upper limb, cardiac, and central nervous system defects. In our patient, the anal opening and external genitalia were absent, and baby also had a single umbilical artery. Radiograph of the baby showed absent radius in the right upper limb. However, organ anomalies could not be ruled out as autopsy was not done.

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