A rare case of heterophagus twinning

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Introduction
The accessory limb or heterophagus twinning is a rare anomaly. There are only a few recorded cases worldwide. Often they accompany spina bifida or other anomalies of spinal structures. Hypothetical theories have been described, including a defect in the early genesis of limbs associated with a neural tube defect, a consequence of a parasitic twin and the presence of a teratoma. We report a child delivered with a well-developed skeletal component with no muscular component arising from the spine.

Case report
A 26 year old mother was admitted to Teaching Hospital, Kandy with spontaneous rupture of membranes at 36 weeks of gestation in her second pregnancy. Up to the current gestation, her antenatal period was uneventful. She has a healthy living child. Dribbling was recorded for 5 hours and the obstetric team decided to let her proceed with normal labour. The mother and the baby were regularly monitored and at delivery the presentation was a foot. Due to obstructed labour, the obstetric team decided to deliver the baby by lower segment caesarean section (LSCS). After delivery, his condition was well. At birth, the following problems were identified. He was small for gestational age, had a large abnormal extension which resembled a lower limb with joint, fingers and an inside bone, had a meningomyelocoele, and had bilateral talipes equinovarus. There were bilateral pre-auricular tags and an umbilical hernia (Figure 1). He was well with normal respiratory and cardiovascular parameters. His haematology was normal with negative blood culture. Ultrasound scans of brain and kidneys were normal.

Hip scan revealed physiological immaturity of both hips. Magnetic resonance imaging (MRI) was done and the findings were: open spinal dysraphism of lower thoracic and upper lumbar vertebrae, meningomyelocoele with dural sac at thoracolumbar junction with protrusion of neural tissue, central canal dilated from cervico-medullary junction to distal end of the cord tethering to meningomyelocoele, an accessory limb attached to the meningomyelocoele sac and consisting of a long bone which could be the femur and a rudimentary foot; this limb was composed of a lipomatous mass covered by skin, with no muscle being visualised; there was no Chiari malformation and ventricles were normal with no hydrocephalus, with other brain findings being compatible with age (Figures 2 and 3). The ultrasound and CT scan findings were compatible with the MRI findings.

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Figure 1: Baby with umbilical hernia and abnormal extension resembling lower limb

*Permission given by parents to publish photograph

On electromyography (EMG) there were no detectable functional motor units in this accessory limb. The findings of the EMG were normal in both lower limbs.

He underwent neurosurgical intervention on the 22nd day of life and the immediate post-surgical period was uneventful. However, during successive days he had CSF leak and lower limb weakness was noted. On post-surgical day 8, baby passed away possibly due to secondary infection by a virulent organism which did not grow in culture. Histopathology of the resected limb revealed that the anatomy of the accessory limb was normal with appropriate histology and no abnormal tissues components were seen with no neurovascular bundle being noticed.

**Discussion**

At present, there are only a few cases reported with varying degree of defect and associations\(^1\)-\(^5\). There is no validated evidence of categorisation or causation including prognostication of this rare entity of anomaly\(^1\)-\(^3\). Different nomenclatures are used. Some of them are: Heterotopic redundancy, aborted twinning, teratoma, tripedus, disorganisation like syndrome, heterophagus twinning, spinal hamartoma, mature teratoma, dorsal accessory limb, midline dorsal appendages, poor parasitic twin etc.\(^1\)-\(^5\). One theory describes it occurring as a result of spinal dysraphism\(^1\)-\(^5\). However, according to the latest hypothesis it follows the primary and secondary neural tube defect leading to spina bifida\(^1\)-\(^2\). As a sequence, primary neural tube defect leads to spina bifida. The secondary neural tube defect leads to over secretion of neural tube fluid which then differentiate to lipomatous, cartilaginous, skeletal, muscular and neural structure developing an accessory limb\(^1\)-\(^2\). Some authors find that heterophagus is due to the development of an asymmetrical twin\(^2\). One twin monopolizes the placental blood supply at the expense of other twin. Consequently, there is ischaemic atrophy of the latter. The remaining part of this developmental anomaly grows as a parasite. In some literature it is accepted that the theory behind this phenomenon is fusion hypothesis in embryogenesis\(^2\).

In this case, the limb was attached at the thoracolumbar region with full phenotypic lower limb and the histology excludes a teratoma. As the spine is mesodermal in origin there can be several structures of mesodermal origin. It supports the presence of spina bifida and meningomyelocele of the same origin.
References

1. Murphy RF, Cohen BH, Muhlbauer MS, Eubanks JW, Sawyer JR, Moisan A, et al. An accessory limb with lipo-
https://doi.org/10.1007/s00383-013-3269-9
PMid: 23392915


https://doi.org/10.4174/astr.2014.87.4.213
PMid: 25317418 PMCID: PMC4196430


https://doi.org/10.1007/s003830050568
PMid: 10370036