Case report
A 7-month-old boy was found to have developmental delay, abnormal muscle tone, and abnormal eye movement in December 2012. Physical examination of the eyes revealed wandering gaze with convergent squint. Ophthalmology was consulted and bilateral retrolental masses were suspected. Blood tests revealed elevated serum creatine kinase level. Computed tomography of orbits showed bilateral hyperdense retrolental tubular opacities with small retinal haemorrhage on the right (Fig 1). Computed tomography of brain also showed communicating hydrocephalus. Magnetic resonance imaging of orbits showed deformed bilateral globes, abnormal T1-weighted and T2-weighted hypo-to-iso-intense contrast-enhancing triangular bands with base near the optic disc and apex at the posterior surface of lens, compatible with bilateral persistent hyperplastic primary vitreous. T1- and T2-weighted hyperintensity at right vitreous body was compatible with previous haemorrhage. Magnetic resonance imaging of the brain showed pachygyria, hydrocephalus, absent septum pellucidum, and hypoplasia of corpus callosum forming type II lissencephaly (Fig 2). Mega cisterna magna, hypoplastic pons, and cerebellar vermis were compatible with posterior cranial fossa malformation (Fig 3). Band-like structures in the bilateral periventricular regions with signal changes similar to grey matter were suggestive of band heterotopic grey matter.

Radiological findings of type II lissencephaly, posterior fossa malformation and retinal anomaly, together with clinical findings of developmental delay, abnormal muscle tone, and elevated serum creatine kinase level were compatible with diagnosis of Walker-Warburg syndrome.

Discussion
Congenital muscular dystrophy (CMD) comprises a heterogeneous group of disorders. Walker-Warburg syndrome is one phenotype of CMD known to occur due to dystroglycanopathy, which is an autosomal recessive condition. The overall incidence is unknown but a survey in Northeastern Italy has reported an incidence rate of 1.2 per 100 000 live births. Walker-Warburg syndrome affects the brain, eye, and muscles with characteristic malformation. Diagnostic criteria for Walker-Warburg syndrome include type II lissencephaly, cerebellar malformation, retinal malformation,
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cleft lip and palate, congenital macrocephaly or microcephaly, posterior encephalocoele, ocular colobomas, congenital cataracts, and genital abnormalities. Neuroimaging findings other than lissencephaly include band heterotopia, cerebellar vermian hypoplasia, dysgenesis of corpus callosum, abnormal white matter changes, hypoplastic cerebral peduncles, intraventricular hemorrhage, cerebellar polymicrogyria, collicular fusion, and fusion of occipital poles.4 Laboratory investigations usually show elevated serum creatine kinase level, myopathic/dystrophic muscle pathology, and altered alpha-dystroglycan.2

Differentiation of Walker-Warburg syndrome from other dystroglycanopathies, for example, muscle-eye-brain disease or Fukuyama CMD, depends on the severity of clinical presentation including motor function and intellectual disability, and involvement of the central nervous system and eye.1 Walker-Warburg syndrome is believed to be the most severe form of dystroglycanopathy with most children dying before the age of 3 years.2 No specific treatment is available for this syndrome. Management is mainly supportive and preventive.

CY Lee *, FRCR, MB, ChB
Department of Radiology, Tuen Mun Hospital, Tuen Mun, Hong Kong

* Corresponding author: prodigycat@gmail.com

References