Congenital hydrocephalus secondary to Walker-Warburg syndrome identified on the Manitoba Neonatal Screening Programme for Duchenne Muscular Dystrophy


Abstract

This report describes our first experience with a clinically important true false positive neonatal screening test for Duchenne muscular dystrophy (DMD). Neonatal screening for DMD began as a pilot programme in Manitoba on 1 January 1986 by analysis of creatine kinase (CK) activity in dried filter paper blood spots. To date, all except two males with positive initial and follow up neonatal CK screening tests were subsequently diagnosed as having DMD. Of these two, one was a newborn male with congenital hydrocephalus whose positive DMD screening test led to the identification of an associated congenital myopathy and confirmation of the diagnosis of Walker-Warburg syndrome.

Between 1 January 1986 and 31 August 1990, 43,513 newborn males have been screened for Duchenne muscular dystrophy by analysis of creatine kinase (CK) activity in dried filter paper blood spots as part of the Manitoba Pilot Neonatal Screening Programme for Duchenne Muscular Dystrophy (DMD) and Becker Muscular Dystrophy (BMD). Eight asymptomatic infant boys have been identified to date with persistently raised CK values and were subsequently diagnosed as having presumptomatic DMD.1 Follow up specimens have been received to date on an additional 35 males who had initial positive neonatal CK screening tests. In all but two the CK levels had become normal at the time of repeat sampling. Of these two, one male was shown to have persistent but benign raised CK-BB isoenzyme.2 We now report a newborn male with congenital hydrocephalus and cataracts who had persistently high CK levels. The positive neonatal screening test for DMD in this child led to the identification of an associated congenital myopathy and subsequent confirmation of the diagnosis of Walker-Warburg syndrome.

Case report

A male infant was born to a 21 year old, healthy, G3P2 mother of Inuit and Caucasian ancestry by caesarean section at 41 weeks' gestation because of prenatal identification of congenital hydrocephalus. The identity of the biological father was uncertain. The pregnancy was complicated by a chlamydial infection at 7 months' gestation and the occasional use of marijuana and hashish. Her first pregnancy had resulted in the birth of a healthy daughter and the second pregnancy ended in an early spontaneous miscarriage. One maternal aunt of the proband was known to have Prader-Willi syndrome with a de novo cytogenetic deletion in 15q11–q13. The proband's maternal first cousin died in infancy from complications of well documented neonatal adrenoleucodystrophy.

On examination after birth, gross macrocrania secondary to congenital hydrocephalus was evident with a head circumference of 48 cm, birth weight 4425 g, and birth length 55 cm. No congenital ocular, facial, musculoskeletal, genitourinary, or internal malformations were immediately evident after birth, but cataracts were noted on the second day of life. The fundi could not be visualised. Neurologically the baby was initially very hypotonic with very poor suck and gag reflexes, but he did not require ventilatory assistance. A right ventriculoperitoneal shunt was inserted on the second day of life. He subsequently tolerated tube feedings, responded to noises, cried loudly, and moved spontaneously. His clinical course was marked by repeated cerebrospinal fluid infections necessitating two shunt revisions. He died at 8 months of age. Permission for necropsy was not obtained. No meaningful social or motor development had been evident.

Chromosome analysis showed a normal male karyotype, 46,XY. TORCH screen was negative. The Pilot Neonatal Screening Programme for Duchenne Muscular Dystrophy reported a grossly raised creatine kinase (CK) level using a modified fluorescent assay performed on filter paper blood spots collected in the routine Manitoba Perinatal Screening Programme as previously described.2 A repeat filter paper assay showed a persistently raised CK level with CK-MM being the predominant isozyme. Two venous CK levels performed at 3 months of age were 3500 and 5368 U/l respectively (normal < 180 U/l). Electromyographic (EMG) examination was consistent with a myopathic process. Ultrasound and CT scan of the head obtained post shunt showed marked dilatation of both lateral, third, and fourth ventricles. Lack of surface gyral markings favoured the diagnosis of lissencephaly (fig 1) and posterior fossa images suggested
cerebellar hypoplasia. Ultrasound of the orbits showed a nodular density in the left orbit anterior to the optic nerve consistent with the presence of a left retinal detachment. A muscle biopsy was obtained from his left quadriceps muscle at 3½ months of age and showed evidence of an active dystrophic process with excessive variation in fibre size, fibre necrosis, and regeneration as well as focal fat replacement (fig 2). Western blot analysis of muscle dystrophin using sheep antidystrophin antiserum (courtesy of Dr L Kunkel and co-workers) and mouse monoclonal antidystrophin antibodies to the rod portion of dystrophin (NCL-DYS1, Novacastra Laboratories, Newcastle) (fig 3) showed abundant dystrophin expression. Immunohistochemical analysis of dystrophin expression confirmed abundant dystrophin staining (data not shown). No detectable alterations were evident in the dystrophin gene using polymerase chain reaction multiplex amplification of eight exons and Southern blot analysis using six separate 32P labelled cDNAs from the American Type Tissue Collection (ATTC, Rockville, Md) that identify over 60 exons spanning the 2 megabase dystrophin gene.

**Discussion**

We propose that our patient had the Walker-Warburg syndrome (WWS) (MIM reference 236670), an autosomal recessive syndrome characterised by severe brain and eye malformations. Type II lissencephaly, cerebellar malformation, and retinal malformation have been present in all reported cases but recently the Walker-Warburg phenotype has been expanded to include congenital muscular dystrophy as a major diagnostic criterion. Documented muscle changes vary widely but all reports include variability of fibre size, central nuclei, fibre splitting, necrosis, and fat infiltration consistent with congenital muscular dystrophy. Although clinical similarities exist with Fukuyama congenital muscular dystrophy, such severe eye abnormalities are uncommon in the latter disorder and it is rarely diagnosed outside Japan. Necessary diagnostic criteria have included type II lissencephaly, cerebellar malformation, and retinal malformation. Raised serum CK has been noted in virtually all patients reported, but the rise in CK varies greatly both between patients (3 to 60 times the predicted normal values) and over time in any individual patient. The identification on a neonatal screening programme of raised CK in this patient led to the assay of the venous CK, the EMG examination, and ultimately to his muscle biopsy which showed active dystrophic pathological changes. The positive identification of congenital muscular dystrophy allowed us to confirm the clinical diagnosis of Walker-Warburg syndrome in this child whose other findings including congenital hydrocephalus, and strong suspicion of type II lissencephaly, retinal malformation, and congenital cataracts by themselves were
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highly suggestive of the Walker-Warburg syndrome. The underlying cause and pathogenesis of the dystrophic muscle seen in Walker-Warburg syndrome is unknown, but we clearly found in this report that it is unrelated to dystrophin deficiency. Whether other dystrophin-like gene products are implicated remains to be determined.

Our pilot neonatal screening programme for DMD/BMD is designed to test the hypothesis that very early identification of DMD boys and subsequent identification of carrier women in their families will lead to a decreased population incidence of DMD and BMD. Approximately 1 in 1000 children have initial positive CK screening tests that require repeat sampling. One child to date has been identified as having persistently raised CK-BB, a benign red cell abnormality. Persistently raised CK-MM on repeat filter paper assay had to date only been seen in our eight true DMD/BMD positives.1 The ascertainment of this child with WWS represents a true false positive. It is thus important to note that other forms of congenital myopathic disorders, including WWS, may be identified on neonatal screening programmes for DMD and BMD. No other male with WWS has been identified clinically between the time of beginning neonatal DMD screening and the ascertainment of this child. In that time period around 50,000 males have been screened and an additional two females with WWS have been diagnosed. This ascertainment of three cases of WWS in 100,000 livebirths suggests a minimal population frequency of ~1/33,333 for this disorder. Since the institution of the Manitoba pilot screening programme, four other males known to us presented in the neonatal period and were subsequently diagnosed as having congenital myopathic disorders (one idiopathic congenital muscular dystrophy, one X linked myotubular myopathy, one phosphofructokinase deficiency, one minicore myopathy). These boys did not have positive neonatal CK screening tests presumably reflecting their only minimally raised venous CK. Nonetheless, we conclude that in addition to true cases of DMD and BMD being ascertained on our neonatal screening programme, other forms of congenital muscular dystrophy may be identified and should be considered when persistently raised CK is detected.

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