Familial olivopontocerebellar atrophy with neonatal onset: a recessively inherited syndrome with systemic and biochemical abnormalities

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SUMMARY Clinical and pathological findings are reported in two siblings who presented in the neonatal period with failure to thrive, hypotonia, pericardial effusions, limitation of joint movement, retinal dystrophy and loss of visual function. Additional features were biochemical evidence of purine overproduction and liver dysfunction. Post mortem, the neuropathological findings in both children were typical of olivopontocerebellar atrophy. It is suggested that the cases represent a recessively inherited inborn error of metabolism.

Definitive diagnosis of olivopontocerebellar atrophy (OPCA) is morphological, requiring the demonstration of degeneration in the cerebellar cortex and its afferent pathways, in contrast to the dentate nucleus and efferent pathways which are usually spared. Since the original descriptions of sporadic1 and dominantly inherited3 forms, a large number of somewhat heterogeneous examples of OPCA have been reported. In an attempt to classify these cases on clinical, pathological and genetic grounds, Konigsmark & Weiner3 divided OPCA into five subtypes, all autosomal dominant except type II cases which were recessive or sporadic. Recently, a putative biochemical defect has been identified in some patients with recessive or sporadic OPCA, deficiency of the enzyme glutamate dehydrogenase4 which is involved in the metabolism of the excitatory neurotransmitter glutamate. Other neurotransmitter abnormalities have been described in dominant OPCA.6

OPCA is rare in childhood, and very few present in the first year of life. To date, all the childhood cases have been part of dominant pedigrees.7 10 We have had the opportunity to examine two siblings presenting a novel and distinctive syndrome with presumed autosomal recessive inheritance, in which OPCA is combined with systemic and biochemical abnormalities.

Case reports

Case 1 was the first child of unrelated Caucasian parents. She was born at term, by normal delivery, weighing 2.8 kg. Apgar scores were satisfactory but after 48 hours she was transferred to the special care baby unit with lethargy, hypothermia and feeding problems. Clinical suspicions of septicaemia were not confirmed by screening but she continued to show failure to thrive. There was obvious oedema of the lower limbs and hepatomegaly, and a pericardial effusion was detected by 2D-Echo when she was 5 weeks old. Weight gain remained poor, diarrhoea was an additional problem, and protein losing enteropathy was suspected. She required several plasma transfusions because of oedema. There was also concern about her vision, when roving eye movements were observed. At the age of 6 months, she was transferred to the Hospital for Sick Children, for investigation of her failure to thrive. Her weight was 3.3 kg, length 55 cm, head circumference 38.5 cm; all measurements were well below the 3rd centile. She was slow to feed, wasted, lethargic, generally hypotonic and developmentally retarded. There were no dysmorphic features. She had occasional roving nystagmoid eye movements, but fixed well and could follow light. Ophthalmoscopic examination showed pale but not hypoplastic discs and pale fundi, diagnosed as retinal dystrophy. The liver edge was palpable 2–3 cm below the costal margin. Limitation of hip abduction was also noted.

Chest radiographs confirmed the presence of a small pericardial effusion, and abdominal ultrasound demonstrated hepatomegaly and ascites. The EEG was normal. The ERGs had abnormally small amplitude and unusual waveforms,
while the VERs were also abnormal. CT scan showed prominent occipital and temporal horns and widening of subarachnoid spaces over the occipital region and medial parietal sulci. The vallecula was also prominent. Biochemical investigations revealed low levels of serum albumin and raised transaminases indicative of liver dysfunction. Subsequently she remained lethargic, requiring tube feeding. Plasma proteins dropped, so she needed plasma transfusions. Her condition rapidly deteriorated with increasing oedema and radiological evidence of pneumonia. Ascites was pronounced and she developed increasing tachypnoea. The diarrhoea continued, but jejunal biopsy and $^{51}$Cr labelled albumin excretion test were normal. She died aged 7 months.

Case 2 The second child, a male, was also born at term after an uneventful pregnancy and delivery, weighing 2-9 kg. Poor feeding and slow weight gain were noted from birth and on the 11th day he was admitted to hospital with recurrent vomiting and diarrhoea. He was referred to the Hospital for Sick Children at 3 weeks: his weight was 2-6 kg, head circumference 35 cm. There were no dysmorphic features, although his appearance was remarkably similar to that of his sister, with rather widely spaced nipples, long fingers and toes, a depressed nasal bridge and slightly long philtrum. Mild flexion contractures of the knees had developed and hip abduction was limited. Mild hepatomegaly was noted. Neurological examination was normal except that he did not fix or follow. Fundal examination revealed pale fundi and an abnormal greyish appearance to the optic discs.

Biochemical abnormalities were similar to the sister as were EEG, VER and ERG. At 10 weeks of age 2D-Echo demonstrated a moderate pericardial effusion and normal intracardiac anatomy. CT scan demonstrated a large vallecula and fourth ventricle with prominent posterior fossa cisterns. Chromosome analysis was normal. On review at 8 months his weight was 4-32 kg, length 61 cm, and head circumference 41-5 cm; all these measurements were below the 3rd centile. There was convergent strabismus, no fixing or following, and the optic discs appeared hypoplastic. Tone was increased in all limbs: the hips could not be fully abducted or knees fully extended. Developmental progress was clearly delayed, but slow rather than regressive. Weight gain remained slow and he died aged 23 months from septicaemia.

Biochemical investigations
Similar abnormalities were noted in both children. Transaminases were raised, indicative of liver dysfunction. Albumin levels were low in Case 1, but not in Case 2. Both siblings had low levels of thyroxin (3-3 and 3-1 nmol/l), yet TSH and free T4 were normal. Thyroid binding globulin levels were low, as were caeruloplasmin levels. Plasma aminoacids, ammonia and urinary organic acids were normal. Levels of plasma urate were consistently elevated in

Fig 1 Case 1 (Age at death 7 months) on the left. Case 2 (Age at death 23 months) on the right. Basal view of the fixed brains. Note the particularly severe shrinkage of the cerebellum and the prominence of the folia, as well as the flattening of the ventral pons. (Parts of the right frontal lobes and cerebellum in Case 2 have been removed prior to fixation).
were unremarkable. Portal necropsy, total tubules, biopsy was symmetrical small and compatible with
acid/creatinine ratio in the urine (2-8) suggestive of purine overproduction. Activity of hypoxanthine-guanine-
phosphoribosyl transferase (HGPRT) was in the normal range (110 μmol/ng/Hb: normal 8–130), but levels of PP
ribose P synthetase were high, 95 nmol/mg/Hb (normal 40–90). Levels of hypoxanthine (100 μmol/l) and xanthine
(31 μmol/l) were also elevated supporting the concept of a defect associated with purine overproduction.

Further investigation in Case 2 demonstrated a high uric acid/creatinine ratio in the urine (2-8) suggestive of purine
overproduction. Activity of hypoxanthine-guanine-phosphoribosyl transferase (HGPRT) was in the normal range (110 μmol/ng/Hb: normal 8–130), but levels of PP ribose P synthetase were high, 95 nmol/mg/Hb (normal 40–90). Levels of hypoxanthine (100 μmol/l) and xanthine (31 μmol/l) were also elevated supporting the concept of a defect associated with purine overproduction.

Plasma bile salts were analysed in Case 2: levels of chenodeoxycholic acid (46-63 μmol/l (N = 0-22 – 12-4) and cholic
acid (17-44 μmol/l (N = 0-06 – 4-55) were raised, compatible with liver dysfunction. Trihydroxyxoprostanic acid
was not detected. Glutamate metabolism was investigated in two ways. Plasma amino acids were analysed following a
standard oral protein load (1g/kg). Methionine levels (1086 μmol/l) were elevated during the initial protein load
but normal in a repeat test. Plasma levels of ketoglutarate were normal during both tests. Also, normal activity of glu-
tamate dehydrogenase was demonstrated histochemically on liver biopsy (Prof B D Lake).

Pathological findings
At necropsy, both children had evidence of marked hepatic portal fibrosis and cystic dilatation of the distal renal
tubules, pulmonary oedema and pericardial effusions. The cardiovascular system was morphologically unremarkable.
Neuropathological findings were similar in both cases and will be described together. Both brains were reduced in size
and weight, and the hindbrains were relatively more affected. Case 1: total weight 510 g (normal for age 690 g):
hindbrain 27 g = 5% of total weight (normal 9%). Case 2: total weight 910 g (normal 1060 g): hindbrain 335 g = 4% of
total weight (normal 12%). The cerebral hemispheres were symmetrical with a normal gyral pattern, and on sectioning
were unremarkable. The cerebellum in each case was very small and shrunken with prominent hard folia, and the pons
was also small with flattening of the base (fig 1). On sectioning the cerebellar folia were thin and white, and the dentate
nuclei difficult to define. The basis pontis was shallow and very pale, while in the medulla the region of the olives was
firm and white.

Histologically, the cerebellar cortex in both cases was dev-
astated. Loss of Purkinje and granule cells (fig 2) was almost
total except in the nodulus in both brains where a few Pur-
kirke cells with abnormal dendritic expansions remained (fig
3). The residual cerebellar cortex was thin, severely gliotic
and virtually devoid of axons. There was no evidence of an
external granule layer in Case 1 as would normally be ex-
pected at 7 months of age. The medullary white matter
was severely depleted of myelin and axons: the dentate ami-
cula appeared worst affected while the hila were better pre-
erved (fig 4). The dentate nuclei themselves showed only
mild cell loss and gliosis (fig 5). The superior cerebellar
peduncles were intact but the fibres of the middle and infe-
tior peduncles were greatly depleted (fig 4). There was severe
neuronal loss and gliosis in the nuclei pontis, virtual dis-
appearance of the transverse pontine fibres (fig 4), and almost
total neuronal loss from the olives with heavy gliosis (fig 6).
Elsewhere in Case 1 there was only very mild cell loss in the
cerebral cortex. Substantia nigra, thalamus and basal gan-
glia, brainstem tegmentum and upper cervical cord were

Neither of the children had any evidence of renal
involvement. In Case 1 there was a small amount of red cell
hemosiderin in the distal renal tubules. In Case 2 there was
a mild patchy interstitial lymphocytic infiltrate in the
kidney, but no other abnormality was detected.

Photoscopic features
Both cases showed prominent gliosis and loss of neuronal
density in the cerebellum. On photography typical features of
Purkinje cell loss were seen in the nodulus (fig 1). There was a
complete loss of Purkinje granule (fig 2) and other nuclear
cells in the cerebellar cortex. This was most prominent in the

Fig 2 Photomicrograph of a folium from the cerebellar
hemisphere. There is complete loss of Purkinje and granule
cells and severe gliosis. (Haematoxylin and eosin × 110).

Fig 3 Cerebellar nodulus. A few granule cells and
degenerate Purkinje cells remain. Note the abnormal Purkinje
dendritic expansion (asteroid body—cactus body). (Arrowhead). (Haematoxylin and eosin × 200).

Gross and Kaltenback's case be classified as OPCA: the cerebellar cortex was hardly affected whereas the dentate nuclei and red nuclei were severely involved. Of other reported cases of childhood onset OPCA, none have been in the neonatal period; the youngest, patient 4 of Colan,7 lost head control and became hypotonic at the age of 5 months. Our cases of OPCA seem therefore to be unique in their early presentation.

Most adult inherited cases of OPCA, and all the childhood cases to date have been part of autosomal dominant pedigrees. In a recent review Koeppen and Barron14 criticise most reports of recessively inherited OPCA from clinical, pathological or genetic standpoint, although they admit that some recessive cases may be incorrectly assumed sporadic for lack of a complete family history. However, in our patients the pattern of inheritance would appear to be autosomal recessive, although there is an important caveat. There are no other children in the family, and both parents are to the best of our knowledge neurologically normal. But they are relatively young; both are still under 35 years of age. In a few families, with autosomal dominant OPCA and retinal degener-

Discussion

The neuropathological findings in both our cases are typical of OPCA. The major damage is sited in the cerebellar cortex and its afferent fibre pathways and nuclei (inferior olives and nuclei pontis), whereas the cerebellar efferent pathway (dentate nucleus and superior cerebellar peduncle) is spared. Morphological changes are remarkably similar in both siblings, except for more generalised neuronal damage in the cerebral hemisphere of the longer survivor. While this might point to non-specific secondary changes, the same basic defect may underlie both cerebellar and cerebral pathology.

OPCA rarely presents in childhood. Berciano11 in a review of 117 reported cases found the average age of onset to be 28 years, and included only two cases12,13 presenting in the neonatal period. However, the morphological changes in the first, Norman and Urich's case, are not those of OPCA, being more akin to pontocerebellar hypoplasia. Nor can the second,
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Fig 6  Horizontal section of the medulla stained for myelin (a) and glial fibres (b). Note the myelin loss and gliosis of the olives and olivo-cerebellar fibres. (a) Luxol fast blue, Cresyl violet × 5; (b), Phosphotungstic acid haematoxylin × 5.

ation, there has been an enormous disparity in age of onset. Childhood onset was within the first 2 years, parental onset in the second or third decade.7 9 15 The most extreme example presented in late infancy, her father and aunt at 41 years and 50 years respectively. These cases7-10 15-17 seem to form a distinct genetic entity from other dominant cerebellar ataxias,18 in particular because of their relatively early onset. But none presented neonatally, or showed the systemic abnormalities which were so evident in our patients.

The clinical features of OPCA are those of progressive cerebellar ataxia, as well as tremor, dysarthria, pyramidal and extrapyramidal signs, ophthalmoplegia and retinal degeneration. In our patients, presenting as neonates, neurological deficit was overshadowed by systemic disease, although limitation of joint movement was detected early in both children, a feature not previously reported in OPCA. Systemic disease, vomiting, diarrhoea, failure to thrive, oedema, pericardial effusions, hepatic fibrosis, has to our knowledge not been recorded before in combination with OPCA. And it clearly points towards an underlying biochemical abnormality.

Some cases of recessive OPCA are reported to have had deficiency of glutamate dehydrogenase.4 5 We were not able to detect any abnormality of glutamic acid metabolism in our patients, although glutamate levels were elevated in one of two protein load tests. The raised levels of methionine, and decreased levels of thyroid binding globulin, caeruloplasmin and albumin could be explained by liver dysfunction and transaminases were elevated. Plasma urate levels were consistently elevated in both siblings, possibly as a consequence of purine overproduction. Spinocerebellar degeneration has been reported in association with a variant form of hypoxanthine guanine phosphoribosyl transferase deficiency,19 but levels of this enzyme were normal in one of our patients. It therefore seems likely that the primary biochemical abnormality in our patients was not directly related to purine metabolism but led to purine overproduction.

So, in conclusion, the remarkable combination in our sibling cases, of neonatal onset OPCA with generalised systemic disease and biochemical abnormalities, is further evidence that inborn errors of metabolism may underlie CNS system degeneration. Current opinion has swung towards that, far from being a specific disease entity, OPCA is a common pathological end point for several clinically and genetically distinct disorders. Our patients appear to demonstrate a new disorder which falls into this group.

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References