Optic nerve hypoplasia, encephalopathy, and neurodevelopmental handicap

John P Burke, Michael O'Keefe, Roger Bowell

Abstract
Abnormalities of the central nervous system are frequently described in optic nerve hypoplasia. In a longitudinal study of 46 consecutive children (32 term, 14 preterm) with bilateral optic nerve hypoplasia 32 (69.5%) had associated neurodevelopmental handicap. Of these, 90% had structural central nervous system abnormalities on computed tomographic brain scans. Neurodevelopmental handicap occurred in 62.5% of the term and 86% of the preterm infants respectively. Term infants had a greater incidence of ventral developmental midline defects and proportionately fewer maternal and/or neonatal complications throughout pregnancy, while encephaloclastic lesions were commoner among the premature infants. An association of optic nerve hypoplasia with the twin transfusion syndrome and prenatal vascular encephalopathies is described.

Optic nerve hypoplasia is a non-progressive congenital abnormality of multifactorial aetiology. Its pathogenesis is not fully understood, but it may follow a variety of insults to any part of the developing visual pathways. It is a major cause of visual loss in childhood and may be associated with important central nervous system and endocrine abnormalities, which include septo-optico dysplasia, porencephaly, encephalolec, congenital suprasellar tumours, hypothyroidism, diabetes insipidus, growth hormone deficiency, and many others. These have recently been the subject of a minireview. The increasing numbers of children described with this abnormality may reflect increased awareness on the part of ophthalmologists as well as an increase in its prevalence.

In contrast to some earlier reports, neurodevelopmental handicaps are being increasingly recognised in association with optic nerve hypoplasia, though their prevalence in and interrelationship with this anomaly have not been fully established. We serially assessed 46 cases of bilateral optic nerve hypoplasia, 32 of which had neurodevelopmental handicap. We examined the factors involved in the simultaneous occurrence of both conditions and describe the association of this anomaly with vascular encephalopathies, in particular the twin transfusion syndrome.

Results
Children with optic nerve hypoplasia and neurodevelopmental handicap were divided into three groups. There were 12 children in group 1 (Table 1) with bilateral optic nerve hypoplasia and neurodevelopmental handicap who were born prematurely and 20 children in group 2 (Table 2) with bilateral optic nerve hypoplasia and neurodevelopmental handicap who were born at or after 37 weeks gestation. There were 14 children in group 3 with bilateral optic nerve hypoplasia who had no neurodevelopmental handicap.

All children in group 1 (Table 1) were born between the 26th and 33rd week, and when last seen their ages ranged from 9 to 56 months. All had been initially referred by paediatricians for visual assessment. Three had been delivered by emergency caesarian section, one for failure to progress in the second stage of labour and two for fetal distress. Four children were products of...
Table 1  Clinical findings in 12 children with bilateral optic nerve hypoplasia and neurodevelopmental handicap born prematurely (≤36 weeks gestation)

<table>
<thead>
<tr>
<th>Patient gestational age (weeks)</th>
<th>Birth history</th>
<th>Pre- and perinatal history</th>
<th>CT or MRI diagnosis</th>
<th>Associated fundus findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>1/28</td>
<td>Assisted breech delivery</td>
<td>Severe RDS GMH-IVH PET and APH</td>
<td>Porencephaly -</td>
<td>-</td>
</tr>
<tr>
<td>2/26</td>
<td>SVD</td>
<td>Severe RDS GMH-IVH</td>
<td>CPVL</td>
<td>Regressed ROP</td>
</tr>
<tr>
<td>3/30</td>
<td>SVD</td>
<td>Suprasellar arachnoid cyst</td>
<td>Atrophic discs -</td>
<td>-</td>
</tr>
<tr>
<td>4/30</td>
<td>SVD</td>
<td>Severe RDS</td>
<td>CPVL</td>
<td>Atrophic discs</td>
</tr>
<tr>
<td>5/32</td>
<td>SVD</td>
<td>Severe RDS</td>
<td>CPVL</td>
<td>-</td>
</tr>
<tr>
<td>6/50</td>
<td>SVD</td>
<td>Severe RDS GMH-IVH</td>
<td>CPVL</td>
<td>-</td>
</tr>
<tr>
<td>7/30</td>
<td>SVD</td>
<td>Severe RDS GMH-IVH</td>
<td>CPVL cerebral atrophy</td>
<td>Regressed ROP</td>
</tr>
<tr>
<td>8/50</td>
<td>SVD</td>
<td>Severe RDS GMH-IVH</td>
<td>CPVL cerebral atrophy</td>
<td>Regressed ROP</td>
</tr>
<tr>
<td>9/30</td>
<td>SVD</td>
<td>Severe RDS GMH-IVH</td>
<td>CPVL cerebral atrophy</td>
<td>Regressed ROP</td>
</tr>
<tr>
<td>10/27</td>
<td>SVD</td>
<td>Severe RDS GMH-IVH</td>
<td>CPVL cerebral atrophy</td>
<td>Regressed ROP</td>
</tr>
<tr>
<td>11/28</td>
<td>SVD</td>
<td>Severe RDS GMH-IVH</td>
<td>CPVL cerebral atrophy</td>
<td>Regressed ROP</td>
</tr>
<tr>
<td>12/27</td>
<td>SVD</td>
<td>Severe RDS GMH-IVH</td>
<td>CPVL cerebral atrophy</td>
<td>Regressed ROP</td>
</tr>
</tbody>
</table>

CT=computed tomographic brain scan. MRI=magnetic resonance imaging scan. RDS=respiratory distress syndrome. GMH-IVH=periventricular leukomalacia. Porencephaly=diffuse cerebral and cerebellar atrophy. APH=antenatal haemorrhage. SLE=systolic lupus erythematosus.

twin pregnancies, and of those suffered from twin transfusion syndrome. Maternal complications included severe pre-eclampsia in two cases, active systemic lupus erythematosus in one, and maternal anaemia of 7-5 g/dl in one mother who initially presented some days before going into spontaneous labour at 30 weeks gestation. There was no history of maternal diabetes mellitus. All 12 infants had a complicated perinatal/neonatal period. Most recent visual acuities ranged from apparently normal to no visual fixation. The amplitude of the binocular flash visual evoked potential correlated better with the visual acuity than did the latent interval. There were six cases with a visual acuity of ≤6/60 (cases 2, 3, 8, 9, 10, 12), three cases with an acuity of ≥6/9 (cases 1, 6, 7), and three cases (4, 5, 11) with intermediate levels. Other ocular abnormalities observed were bilateral optic disc pallor in two cases, presumed cortical visual loss in three, and regressed retinopathy of prematurity in four.

All 12 children had cerebral ultrasound and/or computed tomographic brain (CT) scans performed. Periventricular haemorrhage was visible in six cases by ultrasonography in the neonatal period. One child (case 9) had a normal CT scan. The other 11 scans showed either diffuse cerebral atrophy and/or other encephaloclastic lesions of the developing brain (Table 2). The latter arise from the destruction of previously formed cerebral tissue by a vascular insult. There were no isolated anterior midline developmental defects.

There were 20 children in group 2 (Table 2). When they were last seen their ages ranged from 13 to 80 months. All were referred by paediatricians because of suspected visual handicap. There was no history of maternal diabetes mellitus. Eighteen births were uncomplicated. One mother with twins had an emergency forceps delivery for failure to progress in the second stage of labour (case 21), and one infant had perinatal asphyxia in association with meconium aspiration. Five infants had congenital hydrocephalus and one had cranial diabetes insipidus. An intracranial infection was suspected in three children (cases 16, 24, 26), though unproved in two. The third had cytomegalovirus isolated from her urine three days postnatally (case 24). Eight infants had agenesis of the corpus callosum on CT scan. In addition, three of these had primary and one (case 18) had secondary microcephaly.

Visual acuities ranged from probably normal to no visual fixation. The amplitude of the binocular flash visual evoked potential correlated better with the visual acuities than the latent interval. There were three cases with visual acuities of ≤6/60 (cases 15, 25, 26), one with an acuity of >6/12 (case 20), and the remainder with acuities of ≤6/60. Seven cases had optic disc pallor in association with hypoplastic discs.

There were 14 children with no evidence of neurodevelopmental handicap when assessed (group 3). Three were born before 32 weeks and the remainder after 36 weeks gestation. There was no maternal history of diabetes mellitus or epilepsy. Two of the premature infants were initially referred by paediatricians for retinopathy of prematurity (ROP) screening. One had bilateral optic nerve hypoplasia in association with mild ROP, while the other, who was born to a mother with pre-eclamptic toxaemia, had isolated optic nerve hypoplasia, and thin optic nerves were subsequently identified on CT scan. The third infant was referred at 5 months with early onset esotropia. Of the 11 children born after 36 weeks gestation five were referred by paediatricians with suspected visual handicap. The diagnoses of the referred patients included ocuocutaneous albinism, fetal alcohol effects,
dysmorphic syndrome, postencephalopathic illness, and nystagmus secondary to isolated optic nerve hypoplasia. Six children were referred by their family practitioner. Of these two had albinism, three presented with strabismus, and one had nystagmus, an alternating esotropia, bilateral optic nerve hypoplasia, and a normal CT scan. None of the 46 children studied had tumours of visual pathways.

Discussion
Optic nerve hypoplasia is not a specific disease but a morphological diagnosis which may result from damage at any location in the developing visual pathways. Likewise the location of the injury to the developing brain, causing neurodevelopmental handicap, may vary in accordance with the time in utero when the fetus was exposed to the insult, while its sequelae may or may not be visible on CT scan. This may explain why intrauterine insults sufficient to produce neurodevelopmental handicap may also result in optic nerve hypoplasia.

Skarf and Hoyt noted that delayed development was the most frequently associated nonvisual complication. Nineteen of 41 children (46%) with symmetrical bilateral optic nerve hypoplasia had developmental delay. Margalith et al. noted neuropsychiatric handicaps in 72-6% of 51 patients (92% bilateral), of which mental retardation (70-6%) and cerebral palsy (56-9%) were the most frequent. Edwards and Layden described a series of 14 cases of bilateral optic nerve hypoplasia of which seven had evidence of cerebral palsy and/or mental handicap. Neuroradiological abnormalities were detected in 61% of the 93 patients with bilateral optic nerve hypoplasia in these three series. The results differed from those of earlier studies, as Walton and Robb reported on 20 patients with optic nerve hypoplasia (12 bilateral) who enjoyed normal growth and development, while Abers commented on the infrequent association of optic nerve hypoplasia with mental and/or physical handicap among a series of 45 cases (40 bilateral). In our series 69-5% of 46 cases had neurodevelopmental handicap, and 90% of these had neuroradiological abnormalities. Our database was selective in that 40 of the 46 children were referred by paediatricians for screening or with suspected visual handicap.

It was possible to divide the neurodevelopmentally handicapped children into two groups on the basis of gestational age. There were 12 cases born prematurely and 20 born after 36 weeks gestation. We, like Margalith et al, found the prevalence of perinatal/neonatal and maternal complications to be substantially higher among infants born prematurely. Four of the 12 premature children also had regressed retinopathy of prematurity and germinal matrix-intraventricular haemorrhages. Brown et al. observed a positive correlation between retinopathy of prematurity and intraventricular haemorrhage in premature infants, while others have suggested that severe retinopathy of prematurity and periventricular leucomalacia may both be associated with episodes of cerebrovascular insufficiency. We suggest that the simultaneous occurrence of optic nerve hypoplasia in such patients may imply a contributory ischaemic basis for all three events. To our knowledge no study exists which examines the link, if any, between retinopathy of prematurity and optic nerve hypoplasia.

Ventral developmental midline defects occurred exclusively in the children born at or near term. These represent anomalies in closure of the anterior neural tube, and severe defects may be associated with epilepsy and mental/physical retardation of variable severity. Seven of eight cases in this series with agenesis of the corpus callosum, optic nerve hypoplasia, and neurodevelopmental handicap were products of apparently normal pregnancies. The aetiology of optic nerve hypoplasia in a number of the other 12 cases was unknown (Table 2). One infant (case 24) had intrauterine cytomegalovirus infection. The significance of this association, which was previously reported by Hittner et al., is unclear.

Congenital hydrocephalus has been infrequently described in association with optic nerve hypoplasia. Ghose did not describe any case in a series of 200 patients with hydrocephalus. We noted four five children born at term with congenital hydrocephalus and optic nerve hypoplasia (four obstructive, one communicating). One of these had the Dandy-Walker syndrome, which is frequently associated with other central nervous system abnormalities. One must assume that either optic nerve hypoplasia predated and is unrelated to the hydrocephalus, or that optic nerve hypoplasia can result from this defect, occurring some time prior to the full development of the visual system.

Likewise the simultaneous occurrence of optic nerve hypoplasia and optic atrophy is being recognised with increasing frequency, while the difficulty in differentiating one from the other in some cases may be owing to an insult to the near completely developed visual pathways. The fact that some believe that hypoplastic discs may be pale or normal in colour further confuses diagnostic interpretation in young children. Margalith et al. described 47 patients with bilateral optic nerve hypoplasia, 16% of whom had associated optic atrophy, while nine of 32 patients (28%) in this series had coexisting optic atrophy.

The pathogenesis of encephalopathy and encephaloclastic lesions in the developing brain is diverse and rarely demonstrable. While the role of cerebral ischaemia is still undetermined, the possibility that retinopathy of prematurity may be a manifestation of cerebral ischaemia was suggested by Williams et al. who identified eight of six in the 12 cases born prematurely. Functional disorders of perfusion such as ischaemia from placental abruption, toxoaemia (cases three and nine), or transient vasculitis (case 4) may play a part. Other postulated aetiologies include deficient maternal nutrition (case 8) and anomalies of placental development. To our knowledge the twin transfusion syndrome (cases 3, 5, 21) has not been previously described in association with optic nerve hypoplasia. It was Benirschke who first reported the death of a non-twin optic ataxia, as a potential cause for problems in the surviving twin as a consequence of disseminated intravascular coagulation or from emboli from the
deceased cotwin entering the circulation of the survivor. Melnick\textsuperscript{3} subsequently stated that 1% of monozygotic twin births have severe brain defects as a consequence of the foregoing mechanisms. They and others\textsuperscript{3} described surviving infants with porencephaly and/or hydranencephaly who were usually severely mentally retarded with microcephaly, spastic diplegia, and seizures.

Optic nerve hypoplasia, particularly if bilateral, may be associated with a large variety of ocular and systemic abnormalities. The selection of patients referred to ophthalmic departments can undoubtedly lead to bias. However, the comparison of our findings with other published series allows further insight into the spectrum of optic nerve hypoplasia. Good visual acuity (≥6/9) was present in a minority (6 of 32) of children, while normal CT scans may be found in cases with visual, mental, and physical handicap, indicating the diverse and unpredictable findings that have come to be expected with this anomaly.\textsuperscript{10} We confirmed the general impression that bilateral optic nerve hypoplasia is associated with neurodevelopmental handicap in a considerable percentage of affected term and preterm infants, while the occurrence of optic nerve hypoplasia and hypoxic-ischaemic brain injury in premature infants with or without associated retinopathy of prematurity may suggest common underlying links and merits further study. Similarly, the temporal relationship and significance of coexisting optic nerve hypoplasia and atrophy needs further elucidation.

All children with bilateral optic nerve hypoplasia, and in particular those with impaired vision and suspected neurodevelopmental handicap, should be considered for a CT brain scan and referral to a paediatrician for assessment of their developmental, general, and endocrinological status.

The authors are grateful to Dr Mary King, paediatric neurologist, for her comments.