The nanophthalmic macula

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Abstract

**Aims**—To define an unusual macular appearance found in association with nanophthalmos.

**Methods**—A case review.

**Results**—Seven children (aged 8 months to 17 years) with nanophthalmos were examined. They all exhibited the same clinical findings of an unusual yellow macula appearance with retinal folds and crowded optic discs. Visual electrophysiology performed in four cases was normal.

**Conclusion**—A distinctive yellow macular pigmentation with associated chorioretinal folds and crowded optic discs is present in nanophthalmos. It is proposed that the retinal folds are due to a disparity between scleral and retinal growth while the macula discoloration is due to a congenital abnormality in arrangement or position of the luteal pigment and is not degenerative. Included in this case series is the second case in the literature of nanophthalmos associated with Kenny's syndrome. Inheritance of nanophthalmos appears to be autosomal recessive.

(Neuroophthalmology 1998;82:276–279)

Nanophthalmos is a rare condition consequent upon arrest in the development of the globe in all dimensions. This occurs after the fetal fissure has closed and without other major malformations. Nanophthalmos has sometimes been considered within the spectrum of microphthalmos rather than as a distinct condition despite it exhibiting well defined morphological and clinical characteristics to separate it. Typically, microphthalmic eyes have other structural changes and are frequently associated with systemic abnormalities, whereas in nanophthalmos other ocular or systemic anomalies are generally absent although there are reports of associations with: cryptorchidism, Kenny's syndrome, Hallerman-Streiff-Francois syndrome, and oculodentodigital syndrome.

Different funduscopic appearances associated with nanophthalmos have been described including macular hypoplasia; retinal cysts and pseudopapilloedema; pigmented retinal dystrophy with optic disc pallor and vascular narrowing; macular cysts; and horizontal retinal macular striae in association with geographic areas of pigmentary retinal epithelial disturbance and choroidal thickening.

We describe seven cases of nanophthalmos with an unusual macular appearance, one of which appears to be associated with Kenny's syndrome, and we discuss the mode of inheritance.

Materials and methods

In all seven cases there was extreme hypermetropia, small corneas, shallow anterior chambers, but normal lens size. Fundal examination revealed chorioretinal folds with an unusual appearance at the macula with optic disc crowding (Figs 1–5). For a summary of the findings see Table 1.

In case 6 a genetics referral was made because he was also noted to be of short stature with unusual facies (Fig 6) although he had a normal intellectual level of development. Radiological investigation revealed thickened long bone cortex with stenotic medullary cavities and, although transient hypocalcaemia could not be proved, a presumptive diagnosis of Kenny's syndrome was made.

**Results**

Seven children aged 8 months to 17 years, with findings consistent with nanophthalmos were examined. All were phakic with excessive hypermetropia (+13.25 to +17.25 DS) without other ophthalmic malformations. They all exhibited the same clinical findings of an

![Figure 1](image1.png)

**Figure 1** Fundus photograph of case 1 with a retinal pigment epithelial disturbance evident along with a yellow appearance at macula, retinal folds, and crowded optic discs.

![Figure 2](image2.png)

**Figure 2** Fundus photograph of case 1 with a retinal pigment epithelial disturbance evident along with a yellow appearance at macula, retinal folds, and crowded optic discs.
unusual yellow macula appearance with retinal folds and crowded optic discs. One case had unusual facies and radiological examination revealed thickened long bone cortex consistent with a diagnosis of Kenny’s syndrome.

Visual electrophysiology was available on four patients with the flash electroretinogram (ERG) within normal amplitude limits although 10% less than the laboratory average. Similarly, pattern visual evoked potential (VEP) was within normal latency and amplitude limits though binocular amplitude was an average 20% smaller than the laboratory average. Fluorescein angiograms were not available.

Discussion
Nanophthalmic eyes are typically deep set, associated with narrow palpebral fissures, and are markedly hypermetropic with axial lengths less than 20 mm. They have thickened sclera, small corneal diameters, and general crowding of the anterior chamber with normal sized lenses, making them prone to angle closure glaucoma. Although hypermetropia is the rule, emmetropia or myopia occurs by corneal or lenticular compensation for the short axial length. Other values that have been proposed as diagnostic variables of this condition include ocular volume reduced to two thirds of normal; reduction in ocular size by 15%, and ultrasonic measurements of axial length 2 SD below the mean which is usually considered as 20 mm or less. Ultrasound studies are important, firstly, to look for thickened sclera but also to exclude other ocular abnormalities. It is also helpful in confirming whether the reduction in size is confined to one segment as in anterior or posterior microphthalmos.

A characteristic of nanophthalmos is that the sclera and cornea appear too small compared with the intraocular contents. Abnormal deposits of glycosaminoglycans and elevated levels of fibronectin, which is involved in cellular adhesion, may contribute to the irregular arrangement of collagen bundles found in the sclera of nanophthalmic eyes. This abnormal sclera probably then fails to grow normally. Accumulation of these abnormal products may also thicken the sclera and provide a hydrophilic surface, causing obstruction of the suprachoroidal drainage pathway. This leads to choroidal
congestion, which can damage the retinal pigment epithelial pump causing accumulation of fluid and subsequent choroidal and retinal non-rhegmatogenous detachments, either insidiously or acutely after intraocular surgery. The abnormal growth of the sclera does not affect the lens, which obtains normal volume leading to a high lens/intraocular ratio. A similar disparity in growth between the sclera and retina probably gives rise to the retinal folds as seen in our patients and reported by others.

The high refractive errors leading to ametric amblyopia, associated strabismus leading to strabismic amblyopia, and macular changes are the leading causes of reduced vision in childhood nanophthalmos. In contrast, the leading causes of visual loss in adults with nanophthalmos are glaucoma and the high rate of complications following intraocular surgery. A disparity in growth between the sclera and retina probably gives rise to the retinal folds as seen in our patients and reported by others.

<table>
<thead>
<tr>
<th>Age seen</th>
<th>Cycloplegic refraction</th>
<th>VA</th>
<th>Colour vision</th>
<th>Axial length (mm)</th>
<th>Lens scleral ultrasound</th>
<th>VEP</th>
<th>ERG</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 7 years</td>
<td>R+16.00DS L+14.50/0.75DC</td>
<td>R 6/15</td>
<td>normal</td>
<td>R 15.9</td>
<td>Lens 4.4 mm, Sclera 2 mm R + L</td>
<td>Pattern VEP of normal latency though marginally smaller than average</td>
<td>Normal mixed rod/cone response</td>
</tr>
<tr>
<td>2 5 years</td>
<td>R+14.50DS L+13.25 DS</td>
<td>R 6/15</td>
<td>normal</td>
<td>R 15.2</td>
<td>Lens 4.2 mm, Sclera 1.4 mm R + L</td>
<td>Pattern VEP normal latency slightly small amplitude</td>
<td>Normal mixed rod/cone response</td>
</tr>
<tr>
<td>3 3 years</td>
<td>R+16.00DS L+16DS/0.5DC</td>
<td>R 6/15</td>
<td>normal</td>
<td>R 16.25DS</td>
<td>Lens 4.2 mm, Sclera 2 mm</td>
<td>Pattern VEP of normal latency, marginally smaller than average</td>
<td>Normal mixed rod/cone response</td>
</tr>
<tr>
<td>4 8 months</td>
<td>R+17DS L+15.5DS</td>
<td>R 7/60</td>
<td>normal</td>
<td>R 16.50DS</td>
<td>Lens 4.2 mm, Sclera 2 mm</td>
<td>Pattern VEP of normal latency, amplitudes less than lab average</td>
<td>Normal mixed rod/cone response</td>
</tr>
<tr>
<td>5 5 years</td>
<td>R+16.25DS L+17.0/1.5DC</td>
<td>R 6/15</td>
<td>normal</td>
<td>R 16.50DS</td>
<td>Lens 4.2 mm, Sclera 2 mm</td>
<td>Pattern VEP of normal latency, amplitudes less than lab average</td>
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<tr>
<td>6 8 years</td>
<td>R+14.50DS L+14.50DS</td>
<td>R 6/15</td>
<td>normal</td>
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<td>Pattern VEP of normal latency, amplitudes less than lab average</td>
<td>Normal mixed rod/cone response</td>
</tr>
<tr>
<td>7 3 years</td>
<td>R+17.00DS L+17.00DS</td>
<td>R 6/15</td>
<td>normal</td>
<td>R 16.50DS</td>
<td>Lens 4.2 mm, Sclera 2 mm</td>
<td>Pattern VEP of normal latency, amplitudes less than lab average</td>
<td>Normal mixed rod/cone response</td>
</tr>
</tbody>
</table>

In all cases the intraocular pressure was normal. All cases also exhibited anterior segments with small corneas and shallow anterior chambers but normal appearing lens size giving rise to a very crowded appearance.

The majority of cases of nanophthalmos reported in the literature are sporadic, but some pedigrees of affected families have shown dominant inheritance and some autosomal recessive inheritance. The cases in this series suggest an autosomal recessive trait on the grounds that two family members were affected in the same sibship with normal parents in two separate families. Although no consanguinity can be established in cases 3 and 4 it is probable the parents were related as they were from a small isolated community.

In 1966, Kenny described two patients with a congenital syndrome comprising of dwarfism, thickened long bone cortex, transient hypocalcaemia, and normal intelligence. These and other patients were initially incorrectly reported as myopic whereas they were in fact very hypermetropic. A further report has indicated Kenny’s syndrome in association with nanophthalmos. In one of our patients there was felt to be unusual facies and radiological investigation confirmed thickened long bone cortex, although transient hypocalcaemia could not be demonstrated.
geneticists still felt this was characteristic of Kenny's syndrome. It is important to be aware of this syndrome as the diagnosis may not be suspected and the patient can undergo dangerous electrolyte disturbances, although this was not present in our patient.