Ablepharon macrostomia syndrome

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Abstract
The association of congenital ablepharon with the absence of eyelashes and eyebrows, a wide mouth (macrostomia), and auricular, nasal, genital, and other systemic anomalies has been termed the ablepharon macrostomia syndrome. One such case is reported which illustrates the importance of immediate postnatal ocular management to minimise severe visual loss.

Various rare congenital anomalies of the eyelids are known to be associated with other developmental defects. In the ablepharon macrostomia syndrome (AMS) complete absence of all eyelids, a large mouth, absence of eyebrows and eyelashes, low set abnormal pinnae, irregular anterior nares, flattened malar eminences, dry, coarse and redundant skin, absence of lanugo, webbed fingers, absent or vestigial nipples, ambiguous genitalia, cryptorchidism, and ventral hernia may be found.

We believe only three cases of AMS have been reported, the first two by McCarthy and West and a third by Hornblass and Reifler. We report a further case highlighting the need for urgent eyelid construction.

Case report
The baby was born in July 1986 to a primigravid mother aged 28 years and non-consanguinous father aged 35 years. There was no family history of congenital abnormalities. Pregnancy had been uncomplicated. The following congenital abnormalities were noted at birth; shallow orbits with inadequate eyelids (Fig 1), hypertelorism, large square mouth (Fig 2), protrusion of maxilla, and malformed and low set ears. The skin was wrinkled and hairless; lashes and eyebrows were absent. The genitalia were ambiguous.

Haematological, biochemical, and hormonal profiles and CT scan of the head were all reported as normal. Chromosomal analysis revealed a normal male karyotype.

On the first postnatal day corneal opacities and gross conjunctival chemosis were noted. By the fourth day bilateral corneal vascularisation and central ulcers had developed. Following intensive topical antibiotics and lubricants, bilateral ‘tarsorrhaphies’ were attempted but had broken down within two days. On the tenth day corneal cover was achieved by supplementing the upper eyelids with bilobed temporal scalp rotation flaps (Fig 3). By age 7 weeks the corneal ulcers had healed, leaving vascularised central opacities. The lids were functioning well. No
fundal details could be seen. The flash visually evoked response showed a small P2 and large amplitude P3 at 250 ms. The electroretinogram was normal. His general development at age 15 months was within normal limits for a partially sighted child.

Topical therapy was gradually reduced to liquid paraffin ointment once daily, and by the age of 20 months the corneal opacities had diminished and forced eye closure was complete on both sides. Aged 27 months he underwent right penetrating keratoplasty and bilateral supplementation of the lateral lid regions by means of C-shaped free grafts from the suprACLAVICULAR fossae. This improved lid function, cosmetic appearance, and vision. Left penetrating keratoplasty is also planned.

Discussion
This case and the three cases of AMS reported previously by McCarthy and West1 and by Hornblass and Reifler2 share features which characterise a distinct condition. All cases show bilateral absence of eyelids, eyebrows, and eyelashes, an apparent failure of lateral lip fusion giving rise to the large fish-like mouth, abnormal ears and nose, ventral hernia, absence of lanugo, ambiguous genitalia, and cryptorchidism.

In contrast, a female case described by Barber et al3 had a remarkably similar facial appearance to the reported cases of AMS but there were other notable differences.4 Cesarino et al4 chose to describe their case as lid agenesis-macrostomia-psychomotor retardation-forehead hypertrichosis because a number of features differed from those in AMS.

It had been suggested that AMS is an autosomal recessive condition,5 as ablepharon may occur in association with cryptophthalmos,6 which shares many similar systemic features with AMS.6-9 However, in AMS the periorbital tissues are not fused with the ocular surface, all cases appear to be sporadic, and all chromosomal studies have shown a normal male karyotype.

The ophthalmic management of these neonates begins at birth, with frequent instillation of ocular lubricants and antibiotics. Moisture chambers with plastic wrappings2 or even inferior rectus section with upward rotation of the globe10 may help as temporary measures.

Most techniques described for four-lid construction in the adult involve stages in which total visual occlusion occurs for some weeks,11,12 which in the neonate could induce amblyopia. In our case, once tarsorrhaphies had failed, the upper eyelids were lengthened by making horizontal relieving incisions down to the levators and sliding them down as bipedicile flaps. The defects were filled by means of bilobed transposition flaps from the temporal scalp on either side (Figs 4, 5). The flaps are formed at right angles to each other and then transposed round into the defect. The resultant defect in the first axis is then closed.13 There was sufficient conjunctiva present to obviate the need for mucosal grafts. This technique was favoured because it utilised the excessive and hairless temporal scalp skin14 and the presepal orbitalis muscle was retained at the lid margin.

Postulated mechanisms for AMS are a primary developmental failure, a destruction/absorption process, or some mechanical interference. These would not, however, explain the other associated abnormalities. Neural crest-derived cells contribute to the connective tissue of the eyelids.15 An ectodermal-mesenchymal induction failure could result in ablepharon and the other manifestations of AMS. The thin dry skin lacking in subcutaneous tissue may indicate a primary skin disorder. Perhaps a defect in skin elastic tissue and therefore skin tension during development results in the observed abnormalities (Donnai D, personal communication, 1986). Thus the eyelids may have been prevented from fusing and thereby completing normal development. There are no reports on skin histology in AMS, but periorbital skin biopsies from our case showed no specific abnormality, and collagen studies gave normal findings.

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