Ankyloblepharon Filiforme Adnatum

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Bilateral multiple fibrous tissue bands bridging the lids were identified upon the birth of a Caucasian male born at 40 weeks of gestation following a spontaneous vaginal delivery and uneventful pregnancy. Physical examination including inspection of the anterior segment of the eyes through the fibrous bands did not reveal any abnormal findings. This rare congenital anomaly is known as ankyloblepharon filiforme adnatum (AFA) and can exist as an isolated finding or as part of well-defined syndromes.

AFA describes a single or multiple bands of fibrous tissue joining the upper and lower eyelids either unilaterally or bilaterally. These connecting strands were found to consist of a vascularized central core surrounded by stratified squamous epithelium [1]. The bands consistently arise from the grey line, located between the meibomian gland orifice line and the eyelash line. During normal prenatal development the eyelids remain fused until the fifth month of gestation; in some cases they may not separate completely until as late as the seventh month in utero.

AFA may be associated with iridogoniodygenesis, often complicated by juvenile glaucoma [2]. Systemic associations include trisomy 18 (Edward’s syndrome) [3], Hay-Wells syndrome [4], the popliteal pterygium syndrome (with intracrural webbing of the lower limbs), and CHAND syndrome (acronym for curly hair, ankyloblepharon, nail dysplasia). Other rare associations include hydrocephalus, meningiocele, imperforate anus, bilateral syndactyly, and structural cardiac defects such as patent duc tus arteriosus and ventricular septal defect [5].

Surgery to free these adhesions should be performed as soon as possible to prevent the risk of occlusional amblyopia (amblyopia secondary to reduced visual stimuli). The fibrous bands were severed using a squint hook and scissors and then trimmed at their insertion to the grey line.

The images show the right and the left eye just before the surgery [Figures 1A and 2A] and immediately after [Figures 1B and 2B]. The following day an ocular examination including a dilated fundus examination did not reveal any additional abnormalities.

The prompt recognition and treatment of AFA in the newborn is critical to prevent deep amblyopia and to rule out possible rare systemic abnormalities.

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References