Long-term outcome of strabismus and ptosis surgery in a mother and daughter with congenital fibrosis of extraocular muscles

Dugotrajni ishod hirurškog lečenja strabizma i ptoze zbog kongenitalne fibroze ekstraokularnih mišića kod majke i ćerke

Branislav Stanković*, Milenko Stojković*, Gordana Vlajković†, Ivana Zdravković‡, Dušica Risović§

Clinical Center of Serbia, *Institute of Ophthalmology, †Institute of Anaesthesiology, ‡Institute of Neurosurgery, Clinical Center Zvezdara, §Ophthalmology Clinic, Belgrade

Abstract

Background. Congenital fibrosis of extraocular muscles (CFEOM) is a very rare congenital condition, characterized by variable amounts of restriction of the extraocular muscles, with or without ptosis. The aim of this report was to describe a severe, atypical, exposure-induced corneal stromal lysis in two patients. Case report. A mother and a daughter with a severe CFEOM were presented. The surgery of both extraocular muscles and ptosis led to a fair outcome in mother even 30 years after, and a very good outcome in daughter 4 years after the treatment. Conclusion. Though frequently challenging and disappointing, surgery of extraocular muscles and ptosis in CFEOM can be favorable even in rather severe cases. To the best of our knowledge, the atypical keratolysis we described has not been highlighted in the literature on CFEOM so far.

Key words: blepharoptosis; strabismus; abnormalities; parents; child; ophthalmologic surgical procedures.

Introduction

The congenital fibrosis syndrome is a rare inherited subset of monogenic isolated syndromes of strabism that presented as nonprogressive restrictive ophthalmoplegia, and that includes classic and atypical forms of congenital fibrosis of the extraocular muscles (CFEOM) and Duane syndrome. The syndrome was originally described by Baumgarten 1 in 1840. In 1879 Heuk 2 showed on a postmortem examination an abnormal insertion and replacement of the extraocular muscles with fibrous tissue. It has been traditionally regarded as a primary eye muscle disease. Recent neuropathologic studies have indicated that these disorders may be the result of a primary neuropathy caused by the maldevelopment of oculomotor, trochlear and abducens cranial nerve nuclei, with secondary myopathic changes 3, 4.

CFEOM is, in most cases, an autosomal dominant congenital condition 5 characterized by the variable amounts of restriction of the extraocular muscles, with or without ptosis. CFEOM has been previously divided into several clinical entities: general fibrosis syndrome, congenital fibrosis of the inferior muscle, strabismus fixus, vertical retraction syndrome and congenital unilateral fibrosis. Several types of presentation of the CFEOM within one family have suggested that these are the variants of the same condition 6.

Surgical treatment of patients with CFEOM is challenging 5, frequently disappointing, and most patients appreciate any, even a minimal improvement 7. Binocular vision is...
rarely attained postoperatively, even with a good surgical result, and a repeated surgery is often necessary if a significant amount of residual deviation gradually develops after surgical intervention.

We present the clinical and long-term surgical outcome of a mother and a daughter with a severe CFEOM.

Case report

*Case 1* – a 38-year-old woman with strabismus and bilateral ptosis which had caused a moderate chin elevation (Figure 1), otherwise healthy, was referred for the evaluation of her daughter with a similar appearance she used to have in infancy. The mother had undergone multiple squint and ptosis surgeries 30 years ago. No documentation on the treatment was available. The best (spectacle) corrected visual acuity was 0.6 and 0.1 in the right and in the left eye, respectively. She had rather unusual, mainly horizontal areas, of stromal thinning with a minimal scar formation and a minimal superficial punctate keratopathy, i.e. a form of trophic stromal lysis due to exposure, in the inferior third of both corneas (Figure 2). Surprisingly, these areas of stromal lysis developed without any symptoms of irritation. However, a gradual decrease in visual acuity occurred because of developing astigmatism. An extremely high mixed oblique astigmatism was measured: in the right eye (RE) +2.0 DSph et – 5.0 Dcyl/23º, and in the left eye (LE) +4.0DSph et -8.25 Dcyl/10º, which followed the extent of corneal thinning in favour of the LE. In anomalous head posture (AHP) there was a 20 prism diopters (PD) left exotropia (XT) with 25 PD hypotropia and amblyopia from the childhood. Elevation was absent in addition to the marked reduction of eye movement in both eyes except abduction, with an increasing vertical deviation of vertical side gaze and convergence of attempted upgaze. She refused additional surgery. Artificial tears and safety polycarbonate glasses were recommended.

*Case 2* – a healthy 12-month-old girl with extreme torticollis and both eyes fixed in extreme downward and inward position (Figure 3), had undergone augmented bilateral medial and inferior rectus recession one month after the presentation. A passive duction testing under general anesthesia revealed a severely limited elevation and abduction in the RE and severely limited elevation and moderately limited abduction in the LE. The medial rectus muscle had the course upward to the insertion, suggesting a lower origin in the orbit. Trying to avoid convergence on upgaze we decided to use graded hang back recession: OD 12.0 lower margin /14.5 upper margin; OS 11.0 lower margin /13.5 upper margin, measured from the limbus. For inferior rectus recession we have done “super maximum” hang back recession of 10.0 mm measured from the insertion with recession of conjunctiva and tenon’s capsule. Postoperatively, she developed an immediate large right consecutive XT and small hypetropia. Subsequently, 10 days later she underwent 2.5 mm medial rectus and 2 mm inferior rectus anteposition and bilateral slightly undercorrected brow suspension with polyamide suborbicular pentagon suture according to Fox. Associated high hyperopia was corrected with the appropriate glasses and a part time patching of the LE was instituted. Four years...
after the surgery (Figure 4) she assumed a nearly normal head posture with residual small right exotropia of 5 PD and 7 PD of hypotropia. Elevation was also absent, eye movements limited, particularly adduction and more RE, but the upward convergence seemed unnoticeable. With prolonged treatment with artificial tears (0.3% preservative free HPMC sid and HPMC 2% once at bedtime) after the ptosis surgery her corneas are tolerating the exposure well, with just a mild occasional superficial punctate keratopathy of RE, without the signs of stromal lysis or scar formation in the cornea.

Discussion

The treatment of CFEOM is primarily surgical and aimed at relieving of uncomfortable head position, the reduction of ptosis, and/or the elimination or reduction of significant misalignment of the eyes 5. Unfortunately, nothing could be done to correct the absence of eye movements 10. In general, the less severe the fibrosis, the better the result of surgery 2. Boergen 11 reported a large series of CFEOM patients in which an aggressive surgical approach led to good results in the daughter 4 years after the treatment. An extreme torticollis is certainly the condition that should be alleviated and a treatment not postponed unnecessarily. When considering augmented surgery on medial rectus muscles for children less than 2 years of age, cautious approach must be used even in cases with a large deviation and extreme restrictions. Early anteposition of recessed inferior rectus muscle was probably unnecessary, as even new smaller amount acted as disinsertion, which is, otherwise, a standard approach. Amblyopia can be successfully treated when recognized early. An appropriate optical correction where significant refractive errors are present as well as amblyopia monitoring is needed. To prevent exposure keratopathy and a more serious damage to the cornea, even with mandatory slight undercorrection in brow suspension procedure, close follow-up and treatment with artificial tears and gel or ointment before sleep when needed is obligatory. Although advocated for six months postoperatively it may be a lifelong need in some patients and is sometimes much desired, as exemplified in our case of corneal stromal lysis, despite the apparent lack of symptoms. In patients in whom exposure keratopathy with severe stromal thinning has developed, glasses with polycarbonate lenses to prevent possible perforation of thinned cornea even after a minor blunt trauma to the eye, are recommended.

Despite numerous accounts of postoperative superficial keratopathy and scar formation 5, 6, 10, to the best of our knowledge, our patient is rather unique in respect to the finding of large areas of deep stromal lysis with almost intact epithelium and minimal peripheral scar formation, both in terms of area and density. This being more so because no symptoms apart from gradually decreasing uncorrected visual acuity were reported by the patient. Although this finding is rather unusual according to the literature (though it may be “hidden” within a set of patients in whom scar formation with no further detail was described) it is not entirely surprising in a setting in which scar formation, albeit in a different tissue, is the pathologic basis of clinical manifestations of the disease. It is tempting to regard corneal thinning in our case as a manifestation of the tendency of corneal tissue to form rather hypotrophic scars or no scars at all in response to chronic trophic stromal lysis due to exposure. This seemingly adds to the spectrum of corneal alterations in CFEOM.

REFERENCES


The paper was received on April 3, 2006.