A rare case report of Blepharophimosis syndrome associated with esotropia and nystagmus

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The authors report here a rare case of blepharophimosis syndrome which was associated with esotropia and nystagmus. Blepharophimosis syndrome is a clinical entity characterized by blepharophimosis, ptosis, epicanthus inversus, and telecanthus. In our case, it was also associated with horizontal nystagmus and esotropia. Our patient also had a bilateral mature cataract. It can either be type 1 or type 2 depending on the associated systemic manifestations. The present study describes a case of BPES type 2 with associated nystagmus and esotropia with a bilateral mature cataract.

Keywords: BPES syndrome, Blepharophimosis, Ptosis, Epicanthus inversus, Telecanthus, Nystagmus and esotropia

Introduction

BPES or blepharophimosis, ptosis, epicanthus inversus syndrome is a quartet of clinical anomalies of eyelids specifically characterized by blepharophimosis, ptosis, epicanthus inversus, and telecanthus. The inheritance is autosomal dominant [1] in nature.

Embryological evidence suggests that upper and lower eyelids fuse together by the 8th week of development and separate again by 5th and 7th month [2]. Humans have been found to have abnormal eyelid development, however the molecular events associated with this eyelid development have not yet been fully understood [3,4].
Some progress has now been made to understand the molecular genetics associated with this syndrome [5]. It is predominantly divided into two types: BPES type 1 and BPES type 2; type 1 refers to the classical syndrome complex associated along with premature ovarian failure, whereas type 2 refers to the syndrome complex exclusively limited to the ocular manifestations. Various ocular manifestations may include lacrimal duct anomalies, strabismus, amblyopia, and refractive errors. Other extraocular manifestations may include broad nasal bridge, low set ears, and short philtrum [6]. Diagnosis of BPES is made on clinical presentation, however various genetic aberrations of the gene FOXL2 have been also implicated.

**Genetics:** Blepharophimosis, ptosis, epicanthus inversus syndrome is an autosomal dominant disorder; the causative gene of which is FOXL2, which is localized to the 3q23 chromosome. The genetic aberrations associated are mainly translocations or interstitial deletions of the 3q23 chromosome. The development of the ovary and the eyelid is affected by the FOXL2 gene [7].

**Case Report**

The present study hereby reports a 65-year-old male who reported to the outpatient department of DY Patil hospital Kolhapur with complaints of decreased vision for the last 6 months. The patient gave no history of systemic disease, ocular trauma, or ocular surgeries in the past and allergy to any medications. The patient had drooping of eyelids since birth and there was a family history of similar eyelid complaints in his father, sister, brother, brother’s daughter, and son (Figure 1).

On ocular examination, the resting gaze of the patient had a backward head tilt with arching of eyebrows [Figure 2]. On examination of the lower lids, it was seen that a vertical fold of skin arose from the lower lid that was inserted in the upper lid medially suggestive of epicanthus inversus [Figure 3]. The patient also had increased distance between the medial canthi of around 35 mm which is suggestive of telecanthus [Figure 4]. The patient presented with horizontal nystagmus. The patient also had esotropia of 45° in the left eye. The patient had severe bilateral ptosis with a flat nasal bridge and there was overaction of the frontalis muscle [Figure 5]. The length of the vertical palpebral fissures is around 3 mm and that of horizontal palpebral fissures was 21 mm which is suggestive of blepharophimosis [Figures 6 and 7]. There was an increase in the size of eyelashes (trichomegaly). There was deficient action of the LPS and absent upper eyelid crease. The corneal examination revealed bilateral opacities in lower 1/3rd of the cornea suggestive of healed exposure keratitis. Conjunctival examination in both eyes was normal. Bilateral lenticular examination showed opacities of pearly white color indicating bilateral mature cataract.

The BCVA of the patient was the perception of light and projection of rays in all four quadrants. Lacrimal sac syringing shows patency of both the nasolacrimal duct. The fundus examination shows hazy media due to cataracts. B-scan of both eyes shows no abnormalities in the posterior segment.

![Fig-2: Initial ocular examination of the patient.](image)

![Fig-3: Arrows indicate epicanthal inversus.](image)
BPES syndrome consists of eyelid surgery [9]. In this report, six members of the same family were affected across three generations. In a separate Chinese study, in a family, there were 12 affected individuals across 3 generations [10].

The surgical management of BPES is usually carried out in two stages, the first of which is to perform a medial canthoplasty for the correction of blepharophimosis, epicanthus inversus, and telecanthus and it is usually done between three to five years. The second stage is the correction of ptosis, done about a year later with a brow suspension procedure. The surgical procedure chosen for medial canthoplasty depends on the size of epicanthal folds. In the case of small epicanthal folds, Y-V canthoplasty is performed and if the epicanthal folds are severe, then double Z-plasty is performed [11].

Ptosis correction in adults is done mostly by frontalis muscle flap suspension. In order to have a good cosmetic outcome and improved muscle function, Decock et al preferred the method of super-maximal resection and frontalis suspension [12].

An alternate procedure in which, medial canthoplasty and ptosis correction is performed simultaneously has also been described [13]. Study shows that one-stage correction with standard surgical techniques is not just safe, but also efficient by two retrospectives, interventional studies performed on 21 subjects [14,15].

The causes of abnormal lower eyelid position have now been researched to greater depths and hence has made possible more targeted surgical reconstruction that allows more natural appearance [16].

Another procedure that has been described for blepharophimosis is lateral canthoplasty described by Aginew [17], in which lateral canthotomy and cantholysis is done after which mobilization of the conjunctiva is done and then attached to the skin at the lateral canthus. The lateral fornix is formed by a double arm suture.

Mustarde’s rectangular flap operation is also considered to give better results than the V-Y procedure as described by Verwey [18,19]. In this procedure, paramarginal incisions are made first after which the lids are stretched. This prevents any damage to the canaliculi and also prevents the development of postoperative ectropion. The medial

**Discussion**

Blepharophimosis was first described by Von Ammon in 1841 and Vignes first described BEPS in 1889 after which it has been repeatedly described in literature throughout the globe [8]. Treatment of
Palpebral ligament is shortened and fixed medially to the periosteum with the help of merselene thread.

In our case, as the vision recovery of the patient was important, a bilateral mature cataract first right eye was operated followed by the left eye after 15 days. Following which lid reconstruction was planned in steps. In the first step, it was planned to do medial canthoplasty and in the second step ptosis correction with silicon, the sling was planned. However, the patient has lost follow up.

Reference


