Floppy eyelid syndrome: review

Síndrome da frouxidão palpebral: revisão da literatura

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ABSTRACT
Floppy eyelid syndrome is characterized by the easy eversion of the upper eyelid which occurs spontaneously during the sleep, causing the exposure of the eye surface and chronic papillary conjunctivitis. Its pathogenesis is not totally defined yet, it is usually more frequent in middle-aged, male obese patients and it is associated with systemic disorders such as obstructive sleep apnea, high blood pressure and diabetes. On the occasions which conservative treatment fails, surgical procedures present good results, including surgical techniques which are constantly evolving.

Keywords: Conjunctiva/pathology; Conjunctivitis; Ectropion; Obesity/complications; Hypertension/complications; Diabetes Mellitus/complications; Sleep apnea, obstructive/complications; Eyelid diseases/surgery; Syndrome; Oculomotor Muscles/surgery; Eyelids/surgery; Corneal diseases

INTRODUCTION
Floppy eyelid syndrome (FES), first reported in 1981 by Culbertson and Ostler(1), is recognized by a loose upper lid that readily everts by pulling it upward(1) (Figure1), a soft, rubbery tarsus and Ostler (1), is recognized by a loose upper lid that readily everts by pulling it upward(1) (Figure1), a soft, rubbery tarsus which occurs spontaneously during the sleep, causing the exposure of the eye surface and chronic papillary conjunctivitis. Its pathogenesis is not totally defined yet, it is usually more frequent in middle-aged, male obese patients and it is associated with systemic disorders such as obstructive sleep apnea, high blood pressure and diabetes. Since the initial description of the floppy eyelid syndrome in 1981, several hypotheses have been made for its pathogenesis. Culbertson and Ostler(1), who first pointed out this condition, postulated that spontaneous nocturnal eversion of the upper eyelid would cause events, such as mechanical insult to the conjunctiva, subsequent papillary conjunctivitis and conjunctival keratinization leading to the loss of tarsal elasticity. They made this hypothesis based on the observation that the ocular signs and symptoms are generally more frequent and/or more severe in the eye corresponding to the side on which the patient prefers to sleep. They also suggested an X-chromosome-linked inheritance pattern or hormone influence associated to the problem, since they only found it in men.

Later, other researchers postulated that a genetically predisposition, like genetic abnormalities in collagen, elastin, or both, associated with age and sleeping patterns, would cause the FES(3,10). The obstructive sleep apnea (OSA), often present in patients with FES, lead to the hypothesis that a common underlying connective tissue disorder would explain the syndrome. Redundancy of tissue in the tarsal plate of lateral canthal tendons might contribute to the development of FES, like the excess of oropharyngeal tissues found in OSA cause upper airway obstruction specially when the person is sleeping(11-12).

Inflammation was also pointed out as a contributing factor to FES. Since blepharochalasis and FES share many of the same ocular findings and significant chronic inflammation are encountered in both conditions, a theory of a common inflammatory pathway was

PATHOGENESIS
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made. Inflammation would cause generalized atrophy and attenuation of muscles, tendons and ligaments. Nevertheless, histological studies of the tarsus showed normal structure along with the presence of chronic inflammatory infiltrate\(^{6,11}\).

Hyperglycemia was another condition that was speculated to be involved in the pathogenesis of FES, because glycine is present in high concentrations in collagen and gelatin, and a patient with floppy lids and hyperglycemia was described. The high concentration of glycine in collagen would induce structural changes in the tarsus\(^{9}\). Goldberg et al., however, observed that hyperglycemia is not classically associated with collagen disorders\(^{9}\).

Another theory claims that a combination of local pressure-induced lid ischemia and systemic hypoventilation followed by reperfusion oxidation injury during sleep (sleep apnea) is responsible for changes, such as chronic inflammation, which leads to higher temperature and loss of the normal barrier against water evaporation. This, in turn, would cause additional damage to the skin and this vicious cycle would aggravate lid inflammation, contributing to meibomian gland dysfunction, a condition that might be associated with FES\(^{12}\) and lid floppiness\(^{11}\). The nonspecific ocular surface irritation related with FES might be due to abnormality in tear film dynamics. There is a high correlation between the eye with worse symptoms with more severe floppy lids and the ocular surface evaporation rate. Tear film alterations is prevalent in patients with FES, most of them due to lipid tear deficiency\(^{4}\), which might be caused by abnormalities of meibomian glands\(^{12}\).

Since the main finding in FES is the laxity of the lids, as the name of the syndrome suggests, it was postulated that alterations in collagen would explain this condition. In fact, mutations in the type V collagen genes COL5A1 and COL5A2 were found in patients with ocular phenotype of classic Ehlers-Danlos syndrome (EDS), a disease characterized by laxity and fragility of soft connective tissues. Type V collagen is a quantitatively minor fibril-forming collagen that is present in type I collagen-rich connective tissues such as dermis, tendon and ligament. Mechanisms producing the abnormalities in those tissues were probably associated with altered regulation of collagen fibrillogenesis due to alterations in heterotypic I/V collagen interactions\(^{11}\). Meanwhile, tarsal collagen appeared normal by immunohistochemical studies using Verhoeff’s modified elastin stain suggesting that abnormal elastogenesis might be associated with this syndrome\(^{12}\). The marked decrease in the amount of elastin within the tarsal plate and eyelid skin, however, might be explained by an up-regulation of elastolytic enzymes, particularly matrix metalloproteinases 7 and 9 (MMP-7 and MMP-9), most probably stimulated by repeated mechanical stress, like eye rubbing and sleeping habits. This up-regulation could also be the cause of eyelash misdirection frequently found in FES\(^{10}\). The up-regulation of matrix metalloproteinase (MMP) expression, by the way, might be due to elevated levels of leptin that has been shown to regulate the expression of MMP-9 in a dose-dependent manner. Besides, hyperleptinemia correlates positively with the severity of obstructive sleep apnea (OSA), a condition frequently present in patients with FES. Therefore, hyperleptinemia may be involved in the pathogenesis of both FES and OSA\(^{16}\).

**Corneal abnormality and systemic conditions related with floppy eyelid syndrome**

**Corneal abnormality**

Culbertson and Tseng\(^{13}\), in a retrospective review of 60 patients with floppy eyelid syndrome followed at the Bascom Palmer Eye Institute found a prevalence of seventy-one per cent of patients with clinically significant corneal abnormalities related with FES, that could affect all layers of cornea. Punctate epithelial keratitis was described as the most common corneal disorder, usually diffuse and found only in the involved eye\(^{11}\).

Another corneal abnormality reported by Culbertson and Tseng\(^{13}\) was keratoconus, present in 10% of all patients. FES associated with keratoconus was also described by Parunovic\(^{22}\), Donnenfeld et al\(^{14}\), and Negris\(^{12}\). Corneal endotheliopathy associated with FES was described as a progressive non guttate dystrophy and as a possible Chandler’s variant of the indocorneal endothelial syndrome\(^{16}\). Other corneal signs include subepithelial scarring and deep neovascularization\(^{16}\).

**Obstructive sleep apnea**

Obstructive sleep apnea (OSA) is characterized by intermittent and repeated interruption of airflow during the sleep due to elevated airway resistance. It causes sleep disorder, waking the patient during the night, resulting in excessive daytime somnolence. The patient could present snoring and arterial hypertension. This syndrome is strongly associated with obesity and FES\(^{16}\). The diagnosis is made by polysomnography and the treatment is based in losing weight. Severe cases may be treated by uvulopalatopharyngoplasty and continuous positive airway pressure (CPAP). The first case associating this entities was reported in 1987 by Gornering and Sonneland\(^{16}\). In 1989, Goldberg et al\(^{15}\) described another patient with this condition due to FES and sleep disorders that they called Pickwickian respiratory syndrome. No conclusion between this association had been taken yet.
In 1990, Young et al. reported 3 patients with FES and OSA and proposed a tissue abnormality to correlate these two syndromes. In 1987, Gommering and Sonnelland selected patients with FES to underwent polysomnography and patients with OSA to be examined for ocular abnormalities and measurement of vertical upper lid pull. In a group of 8 patients with FES, all of them had also OSA in different degrees; however, among 20 patients with OSA, only one case of FES was diagnosed. Nevertheless, measurement of vertical upper lid pull was significantly higher in these patients compared with normal population. McNab also showed one patient with the 2 entities, that improved of FES after treatment of OSA (uvulopalatopharyngoplasty and CPAP), with no papillary conjunctivitis and normal lid laxity.

In 2005, McNab analyzed 50 patients with FES and found that 96% of them had symptoms of OSA. Seventeen of these patients were undergone sleep studies and in sixteen (94%) was confirmed the diagnosis of OSA. This result corroborated with his previous study showing that patients with FES and OSA had worse degree of severity, higher number of apneic or hypopnoeic episodes per hour, and lower minimum saturation of arterial oxygen, when compared with patients who had only OSA. It suggests that FES is a severity marker of OSA. In 2006, Karger et al. confirmed McNab findings. They encountered an association with subjectively easy lid eversion and sleep disorders, but a low frequency of FES. According to currently data, most of the patients with FES have OSA; however, the opposite is not true. Patients with FES should undergo polysomnography to evaluate a possible OSA, whose treatment could give a better life quality to the patient.

Several other systemic conditions have already been associated with FES, such as diabetes, ischemic heart disease, hypercholesterolemia, osteoarthritis, asthma, gastroesophageal reflux disease, chronic renal failure, and schizophrenia. Based on the speculative etiologies of FES, many different kinds of clinical and surgical treatments were proposed. The clinical treatment involves losing weight, taping the eyelids during the night, eye shields and topical lubricants. Improvement of FES manifestations by the treatment of OSA with CPAP was seen by McNab.

The surgical techniques changed in the years and are constantly being improved to treat this condition. Gerner and Hughes first described a shortening procedure in a patient with hyperglycemia and floppy eyelids. Dutton had also tried an eyelid shortening procedure, in which a full thickness resection of the lateral one third to one half of the eyelid was performed with primary layered closure. Later, lateral tarsorhaphy was used by Bouchard. Karesh et al. had done a similar procedure to Dutton’s by using a lateral canthal tendon placentation, and lower eyelid horizontal shortening. Despite of the good effectiveness, these procedures do not have a fine aesthetic result. Therefore, Periman and Sires presented a modified technique in which a temporally placed, modified wedge excision is used to achieve horizontal shortening, with 91% of improvement of symptoms in a follow-up of 52 months.

Valenzuela and Sullivan performed, based on the presence of predominately medial upper laxity, a medial upper eyelid horizontal shortening with an upper eyelid skin reduction when excess of skin was present, and some other patients also underwent horizontal lower eyelid shortening additionally or not to the lower eyelid retractor plication. The authors had a great success in 18 months of follow-up, with relief of the symptoms and good aesthetic appearance.

Besides the surgical management is the main and more effectivite treatment for FES, a long-term study involving 71 patients who had undergone the surgery for FES at Moorfields Eye Hospital demonstrates significant recurrence rates varying from 25.6% to 60.6% depending on the procedure used. The data provided by this showed better survival outcomes using the medial canthal and lateral canthal pllication and upper lid lateral tarsal strip procedures in comparison with the full-thickness wedge excision procedure, which was recommended to be avoided.

**REFERENCES**