The Incidence of Dry Eye Disease Related to Long Term Diabetes Mellitus Tip 2

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Abbreviations: DM: Diabetes Mellitus; DES: Dry Eye Syndrome; TUBT: Tear Break Up Time; BVCA: Best Corrected Visual Acuity; SPK: Superficial Punktate Keratitis; MGD: Meibomian Gland Dysfunction; MG: Meibomian Gland

Introduction

Diabetes is a disease that affects the body’s ability to produce or use insulin effectively to control blood sugar (glucose) levels. DM is a chronic metabolic disorder usually genetic [1,2]. The global diabetes mellitus evidence predicts that 6.4 % of the world population are affecting 286 million adults in 2010 by 2030 is predicted to increase about 7.6 % affecting about 439 million of adults [2]. People with diabetes frequently develop dry eye syndrome [3]. It is estimated that over 50% of people with type 2 diabetes have dry eye [4]. Patients will complain of eye irritation, excessive tearing, fatigue, red eye, gritty sensation, increased watering of the eyes, dryness and blurred vision [5,6].

The causes of dry eye are so many, but the main cause is a disturbance of the tear film due to either decreased tear production or excessive evaporation of tears. MGD occurs when the normal function of the MG becomes compromised because of gland blockages that occur over time, resulting in inadequate oil production into the tear [7]. If the MG malfunctions, the lipid layer may be reduced, allowing rapid evaporation of the tear complex [7-9]. Research shows that most cases of dry eye associated with diabetes are caused by insufficient production of tears due to “autonomic neuropathy. The condition is affecting the nerves that control the lacrimal gland secretion damaging the production of tear film that normally produce moisture to keep the eyes well lubricated [10,11]. When the transparent and sensitive cornea is no longer adequately lubricated, the cells of the cornea become damaged and free nerve endings are exposed. The exposure of the nerve endings leads towards symptoms of dry eye.

Furthermore, by keeping blood sugar levels as firmly controlled as possible is the first step in preventing dry eye syndrome at diabetes patients. The high blood glucose HbA1c more than 6.4 % lead to autonomic neuropathy affecting the tear gland but also affects the quality of our tears by increasing the amount of glucose in the tear layers and disrupting their normal chemical composition [12,13]. The tear fluid contains water, lipids, mucin, lactoferrin, lysozyme, lipocalin, immunoglobulins, glucose, urea, sodium potassium, lacritin [14].

Three layers are components of tear film: the aqueous layer, the mucus layer and the lipid layer. Also the goblet cells of conjunctiva plays very important role by producing and secreting mucin lubricating the ocular surface [7].
Insufficient tear production and changes in osmolarity promote high concentrations of proteins within the tears inducing apoptosis of surface epithelium and a vicious cycle of increased expression of inflammatory cytokines from ocular surface [7]. The condition gets even worse by apoptosis and decreased mucin production by goblet cells [15] (Figure 1&2).

**Purpose**

The purpose of the study is to prescribe and analyze the correlation of Dry Eye with diabetes related to duration of primary disease.

**Methods**

A total number of 98 non selective patients, diagnosed with Diabetes Mellitus Tip II, sent for routine examination from Endocrinologist, underwent ocular examinations. 50 male and 45 female, aged from 42-84 years old.

History of disease, duration of diabetes, age, sex, the level of hemoglobinA1c equal or over than 6.5 % (for the past 3 months) was obtained by reviewing the medical records from the Endocrinology Department and direct patient interview. Visual acuity with correction BVCA, OSDI questionaire for classification of DES. Slit-lamp examination, tear meniscus at inferior lid margin, tear film break-up time (BUT) test, corneal sensitivity test, Schirmmer Test to evaluate the tear film secretion from the lacrimal gland, Rose Bengal stain test for the conjunctiva goblet cells malfunction, fluoresceine corneal dye test to evaluate the corneal epithelial involvement. Also tear film measurement performing Schirmer’s test in 5 min time to evaluate the amount of tear film production from Lacrimal Gland, cotton-swab test for the corneal sensitivity damage and peripheral neuropathy. Tear film break up time test measurement was performed using fluorescein strips.

Visual acuity was evaluated by using Snellen’s chart. Slit lamp (Nidek SL-450) examination was performed with particular attention to lid margin, bulbar and tarsal conjunctiva, cornea and conjunctiva to detect any abnormality of meibomian Gland Disfunction and also the presence of Superficial Punktate keratitis or any involvement-reduction of Golbet cells production. In addition the patient complains of tearing photophobia, red eye, itching, foreign body sensation, increased watering of the eyes, dryness and blurred vision was obtained. The duration of primary disease was recorded and divided in three groups of patients: duration onset of DM -5 years, duration of 5-10 years and more than 10 years of duration (Figure 3-5).

**Results**

From our research the results are as follows. 64 patients were diagnosed with Dry Eye Syndrome of varying degrees from mild to very severe according to OSDI questionnaire, 28 male / 36 female. The diagnosis was made by having two or more positive tests performed as mentioned above and patient’s complaints. In addition, at those subjects diagnosed with DES, 5 patients represent pinpoint corneal dye with fluorosceini (SPK), 8 of them where positive to corneal cotton swab test, 16 of them resulted positive to Schirmer test less than 10 mm in 5 min, 14 of them was showing TBUT test less than 9 sec, presence of positive tear meniscus was seen in 15 patients, Rose Bengal staining of conjunctiva was detected in 6 patients, MGD was noticed at 12 subjects. Regarding the duration of Diabetes in our study the
findings where: 21 patients were diagnosed with DM during the period of 5 years of duration, 31 of the patients were diagnosed with DM since 5-10 years ago, 46 of them had Type 2 more than 10 years.

With insight at the group with 5 years of duration there was a low number of patients presenting very mild dry eye at 11 patients, at group with 5-10 years of duration of DM there were about 20 patients presenting mild and severe dry eye, considering that the group included patient with DM more than 10 years of duration; 31 patients were presenting severe to very severe DES (Figure 6 & 7).

Figure 6: Photo Corneal staining SPK.

Figure 7: Photo Rose -Bengal staining of conjunctiva.

Conclusion

This study, the Dry Eye Syndrome showed to have a high correlation with Diabetes Mellitus Tip II (about 56.9%). Prevalence and severity of Dry Eye was significantly higher at patients with longer duration of diabetes (46.9%) and poor glycaemic control, less high at the patients with no longer duration of primary disease. Dry Eye seems to be an important contributing factor related to corneal abnormalities by disrupting the normal chemical composition of the tear film layers. In our study we have shown that the Age and Sex seem not to play an important role in this condition. Good glycaemic control is very important and decisive for prevention of the DES.

Discussion

To conclude, everything depends on blood sugar levels (HbA1c). The higher the attention monitoring the blood sugar level consists in our first step in preventing dry eye syndrome associated with diabetes [12]. More often, the patients are unaware that their blood glucose is out of control, which is fundamental for the treatment at diabetes related to dry eye syndrome [16]. Educating the patients by sharing our knowledge of primary disease DM, the proper treatment, the appropriate monitoring, the diet instructions the exercises and follow up by Endocrinologist is the key to improve the quality of their life style [2] and to reduce the risk for developing DES as much as possible.

Subsequently, it is very important to achieve a good collaboration among physicians to detect early the DM Type 2, to receive the proper management and treatment options for the patient [17]. Compliance of the patients is also a very important factor to learn how to live with diabetes and to present regularly for follow-up. The DM is a chronic disease hence either the DES remain chronic disease as well, for that reason is still very difficult to manage both conditions [15]. New modalities need to be developed though collaborative research and further studies need to be undertaken to treat the DES and the primary disease DM constantly (Figure 8).

| Table 1. Dry eye severity grading scheme |
|------------------------------------------|
| **Dry Eye Severity Level** | **1** | **2** | **3** | **4** |
| Discomfort, severity & frequency | Mild and/or episodic, occurs under environmental stress | Moderate episodic or chronic, stress or no stress | Severe frequent or constant without stress | Severe and/or disabling and constant |
| Visual symptoms | None or episodic mild fatigue | Severe or chronic episodic fatigue | Severe chronic and disabling fatigue |
| Conjunctival injection & irritation | None to mild | None to mild | None to mild | None to mild |
| Conjunctival staining | None to mild | Mild to moderate | Moderate to marked | Marked |
| Corneal staining (severity/size) | None to mild | Variable | Moderate to marked | Marked |
| Corneal/ocular signs | None to mild | Mild debris, corneal vascularization | Flimbrane type stains, capillary loops | Fine capillary loops, capillary loops |
| Lid/lacrimal issues | MGD bizarre present | MGD bizarre present | MGD bizarre present | MGD bizarre present |
| TFBUT (sec) | Variable | ≤ 10 | ≤ 5 | ≤ 2 |
| Schirmer test (mm 5 min) | Variable | ≤ 10 | ≤ 5 | ≤ 2 |

* Most these signs and symptoms. TFBUT: Fluorescein breakup time, MGD: meibomian gland disease.

Figure 8: Photo DES Delphi Panel grading scheme.
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