ESTIMATION OF TOTAL IgE IN THE LACRIMAL FLUID AND SERUM OF PATIENTS WITH GIANT PAPILLARY CONJUNCTIVITIS

By
El-Nafees R., Mokbel Th.,
El-Lakany R., Salvi N.*

From
* Clinical Pathology
Ophtalmology Department, Faculty of Medicine Mansoura University
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INTRODUCTION
Giant papillary conjunctivitis (GPC) is a disease of contact lens wearers, which was first reported by Spring (1974), in 78 out of 170 soft lens wearers. The disease is characterized by the appearance of giant papillae more than 1 mm in diameter at the upper tarsal conjunctiva with subjective symptoms including cloudiness and dislocation of the contact lens, mucoid discharge, foreign body sensation, irritation, watering, hyperaemia, and itching (Allansmith et al, 1977). The disease may occur in both hard and soft contact lens wearers but the physical appearance differs. Papillae are fewer in rigid lens GPC, and more likely have flat tops in contrast to papillae of soft lens GPC which are more rounded (Henriquez et al, 1980). Biopsy specimens of the upper tarsal area showed normal conjunctiva with micropapillae and huge expansion of tissue which is mostly collagen (Korb et al, 1980). Histopathological studies showed the presence of mast cells in the conjunctival epithelium and substantia propria,eosinophils and occasionally basophils in the epithelium or substantia propria (Allansmith et al, 1978).

The etiology of GPC is not completely clear. The possible causes include repeated mechanical irritation of the conjunctiva by the contact lens with lid movements, allergy to the lens

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material, history of atopy, lens deposits, irritation by the cleaning solutions, and poor lens hygiene (Momose et al, 1983).

A specific immunological response to antigenic material present in the coating on the surface of the contact lenses including mucous, protein, cells, cell debris, bacteria, and airborne pollutants may have a role in the development of GPC (Fowler & Allansmith, 1988).

Takeuch et al., (1983) studied the atopic history and the Ig E values in blood for 22 patients with GPC. Atopic history was found in 7 patients and serum Ig E was elevated more than 500 lu/ml in 4 patients. Ig E values in serum and lacrimal fluid in different atopic conjunctivitis patients were found to be higher than in non-atopic conjunctivitis patients and in normal control subjects (Aalders- Deenstra et al, 1985).

SUBJECTS & METHODS
A group of contact lens wearers having GPC and another group of normal contact lens wearers were randomly selected for this study. All were soft lens wearers (high water content, extended wear) Hard lens wearers were not excluded but due to the infrequent use of hard lenses none of them was received during the study.

History:
Full detailed history was taken including history of atopic diseases in the patient or the family, duration of contact lens use, types of lenses used, method of lens care, and the nature of symptoms.

Examination:
Full ocular examination was carried out including slit lamp examination of the anterior segment, conjunctiva, and inspection of the contact lens.

Collection of lacrimal fluid:
The lacrimal fluid was collected using a glass capillary tubes the capacity of which is 100 µl. No surface anaesthetic was used to avoid altering the chemical composition of the lacrimal fluid.
Each glass capillary was put in the lower fornix gently until filled, then transferred to the test tube and another one used to collect more lacrimal fluid. 500-800 μl were collected for each patient. The lacrimal fluid was stored in polystyrene tubes at -70°C till analysis was carried out.

A blood sample was taken for each patient at the laboratory for the estimation of serum IgE.

IgE determination:
The total IgE content of the lacrimal fluid and serum was measured with the commercially available Phadezyn PRIST (Pharmacia - Sweden) according to the instructions given by the manufacturer. The value is expressed in IU/ml.

RESULTS
28 GPC patients were studied. 20 were females and 8 were males. Their ages ranged between 16-40 years. 22 normal contact lens wearers were also studied. Their ages ranged between 18-38 years. All the studied cases used soft contact lenses for a duration ranging between 3-10 years.

Atopic History:
History of atopic diseases was detected in 8 out of the 28 GPC cases and in 2 out of the 22 normal contact lens wearers. Weather in the patient or the family.

Lacrical and Serum IgE:
The results of total lacrimal and serum IgE in the GPC patients and in the normal contact lens wearers are shown in Table I.

Table I, total lacrimal and serum IgE (IU/ml) in GPC patients and normal contact lens wearers.

<table>
<thead>
<tr>
<th></th>
<th>GPC patients</th>
<th>Normal lens wearers</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No.</td>
<td>%</td>
</tr>
<tr>
<td>Normal</td>
<td>4</td>
<td>14.29%</td>
</tr>
<tr>
<td>Elevated</td>
<td>24</td>
<td>85.71%</td>
</tr>
<tr>
<td>Total</td>
<td>28</td>
<td>100.00%</td>
</tr>
</tbody>
</table>

24 (85.71%) out of the 28 GPC patients showed elevated lacrimal IgE levels (5.7 - 63.8) IU/ml and also elevated serum IgE levels (205.6 - 943.7) IU/ml.
The remaining 4 patients showed normal lacrimal IgE levels (0.35 - 0.56) IU/ml as well as normal serum IgE levels (16.4 - 137.6) IU/ml.

Only 3 (13.63%) out of the 22 normal soft contact lens wearers showed elevated lacrimal IgE levels (8.2, 15.3, 23.7) IU/ml and also elevated serum IgE levels (183, 241.5, 341.8) IU/ml. The remaining 19 (86.37%) cases showed both normal lacrimal IgE levels (0.37-0.83) IU/ml, and serum IgE levels (7.6 - 124.8) IU/ml.

**DISCUSSION**

Many factors have been suggested to contribute in the establishment of GPC in contact lens wearers but the exact aetiology is still not completely clear.

The possibility of being an immunological process was raised by Korb et al., (1980) depending upon the nature of cellular infiltration. The study of IgE in serum and in the lacrimal fluid became an important way to diagnose atopic conditions and to explain the occurrence of certain diseases on immunological basis. In our study the total serum and lacrimal IgE were estimated using the PRIST which was found by Ballow & Mendelson (1980) to be able to measure only values above 0.25 IU/ml, and by Liotet et al., (1983) to measure total levels above 0.5 IU/ml. A considerable elevation was detected in both lacrimal and serum total IgE in 85.7% out of 28 GPC cases, with the elevation more pronounced in the lacrimal fluid which may reflect local production of IgE in the lacrimal fluid. A slight elevation of total lacrimal and serum IgE was detected in only 3 out of 22 normal contact lens wearers, 2 of them had +ve atopic history but non of them had conjunctivitis. 8 out of the 28 GPC cases had +ve atopic history together with elevated IgE levels. These results show that the occurrence of GPC is not just a conjunctival response to local irritation of the lens coating, or cleaning solutions, but is rather an immunological process. The local factors can stimulate the development of type I and type IV hypersensitivity responses. Our results agree with those of Takeuchi et al., (1983) who considered
GPC a sort of immediate hypersensitivity reaction.

**SUMMARY**

Total Ig E levels in the lacrimal fluid and serum were estimated in 28 soft contact lens wearers having giant papillary conjunctivitis (GPC) and in 22 normal soft contact lens wearers using the paper radioimmunosorbent test (PRIST).

A considerable elevation of lacrimal and serum IgE was detected in 24 (85.71%) out of the 28 GPC patients. The elevation was more pronounced in lacrimal than in serum IgE.

A slight elevation was detected in only 3 (13.63%) out of the 22 normal contact lens wearers.

**REFERENCES**


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الملخص العربي
تحديد نسبة الأجسام المناعية المضادة (أي) الكلية في السائل الدمعي والدم في مرضى التهاب الملتحمة الحلمي العكالق
تم تحديد نسبة الأجسام المناعية المضادة (أي) الكلية في السائل الدمعي والدم في 28 مريضاً مصابين بالتهاب الملتحمة الحلمي العكالق بلبسون عدسات ملتصقة رخوة، وفي 22 شخصاً طبيعياً بلبسون عدسات ملتصقة رخوة، باستخدام اختبار ورقة المص المناعي المشع.

وقد لوحظ وجود ارتفاع ملحوظ في الأجسام المناعية المضادة (أي) في السائل الدمعي والدم في 24 مريضاً (71.5٪) ضمن 28 مريضاً بالتهاب الملتحمة الحلمي العكالق، وهذا الارتفاع كان أكبر في السائل الدمعي منه في الدم.

كما وجد ارتفاع طفيف في 3 مرضى (12.5٪) ضمن 22 شخصاً طبيعياً بلبسون عدسات ملتصقة رخوة.