Altered hyaluronic acid content in tear fluid of patients with adenoviral conjunctivitis

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ABSTRACT

The adenoviral conjunctivitis is one of the biggest causes of conjunctival infection in the world. Conjunctivitis causes relatively nonspecific symptoms, as hyperaemia and chemosis. Even after biomicroscopy, complex laboratory tests, such as viral culture, are necessary to identify the pathogen or its etiology. To contribute to the better understanding of the pathobiology of the adenoviral conjunctivitis, the tear fluids of patients with unilateral acute adenovirus conjunctivitis (UAAC), normal donors (control) and patients with allergic conjunctivitis were analyzed. Tear samples were collected with Schirmer strips from control, allergic conjunctivitis and UAAC patients, diagnosed by clinical signs. UAAC tears were tested positive in viral cultures. After the elution, HA was quantified using an ELISA-like fluorometric assay and the protein profile was determined by SDS-PAGE. A profound increase in the HA tear content in UAAC patients was found when compared to control and ALC. This HA increase in UAAC tears remarkably was not observed in tears from contralateral eyes without clinical signs, nor in allergic conjunctivitis. In addition a distinct profile of UAAC tear proteins was observed in patients with UAAC. The quantification of HA in the tear fluid is a rapid, sensitive and specific test. This molecule might be a biomarker candidate for acute conjunctivitis.

Key words: conjunctivitis, glycosaminoglycans, hyaluronic acid, tear film.

INTRODUCTION

The adenovirus conjunctivitis is one of the most common conjunctival infection. Normally it does not result in vision loss, but it frequently causes abstention from school or work, with a high economic and social cost (Okada and Forrester 2000, Butt and Chodosh 2006). Although its pathogenesis is not completely understood, the role of extracellular matrix components has recently been studied (Natividad et al. 2006).
A healthy ocular surface requires a functional tear film. The normal function of tear fluid is due to its complex biochemical composition consisting of buffered electrolytes and a diversity of proteins and glycoconjugates (Van Haeringen 1981, Baker et al. 2006).

Hyaluronic acid (HA), an important component of the extracellular matrix, is a large glycosaminoglycan composed of repeating units of β-D-glucuronic acid and N-acetyl-D-glucosamine. HA plays an important role in tissue development, cell migration, cell proliferation and inflammation (Inoue and Katakami 1993, Gomes et al. 2004). HA is increased in the tear fluid when corneal epithelium erosion is present, and may play an important role in corneal epithelium wound healing (Oya et al. 1995, Miyauchi et al. 1996). Tear fluid consistency (gel-like) can be attributed to HA (Itano et al. 1999). Another important characteristic of this GAG is its chemical structure that can attract ions and water due to a negative charge density (Frescura et al. 1994, Yoshida et al. 1996).

Endogenous hyaluronan is present in virtually all corneal disorders (Fitzsimmons et al. 1994). HA is normally detected only in the corneal endothelium, and the presence of any amount of this compound in the epithelium or stroma is likely to indicate altered tissue (Inoue and Katakami 1993). Vitreous humor contains HA and its receptor (CD44), and these molecules are also present at the apical surface of corneal endothelium (Tengblad 1979).

Tear proteins also play an important role in maintaining eye surface integrity and in patients with external eye diseases (Avisar et al. 1981, Van Haeringen 1981, Ballow et al. 1987). The protein fraction of normal tears contains antimicrobial factors which are important for protecting the external eye from infection. These substances are produced by the main and accessory lacrimal glands. Important components of the hosts defense system for the external eye include complement proteins, immunoglobulins, especially secretory IgA (sIgA), lysozyme, lipocalin (TSPA) and lactoferrin (Friedman 1990, Kuizenga et al. 1991, Tragoulias et al. 2005). Lactoferrin, an iron complexing protein in normal tears, has bacteriostatic, bactericidal and complement (C) inhibitory activity properties which make this tear protein an important component of the nonspecific host defense system of the external eye (Ballow et al. 1987, Flanagan and Willcox 2009). Human tear prealbumin, now called tear lipocalin, was described as a major protein in tear fluid, and is the main lipid binding protein in tears. It exerts important functions in eyelid lubrication, and acting as a general protection factor for cornea and conjunctiva epithelia (Redl 2000). IgA is the only significant immunoglobulin found in tear film. The antibodies are postulated to play a role in preventing adherence of microorganisms to the ocular surface (Alizadeh et al. 2001, Knop and Knop 2005). Another component of tears is lysozyme, a bacteriolysic protein first described by Fleming (Fleming 1922, McClellan 1997, Caffery et al. 2008). An allergen exposure can cause a marked increase in lysozyme secretion (Proud et al. 1998).

The aim of this study was to quantify HA content and verify the protein profile of tear fluids from patients with adenoviral conjunctivitis and compare them with normal donors.

MATERIALS AND METHODS

SUBJECTS

Tear samples were obtained from 53 patients; 15 eyes affected by acute adenoviral conjunctivitis, diagnosed from clinical signs, without epithelial defects and testing positive for viral culture and 15 contralateral eyes without clinical signs; as well as both eyes from 23 normal subjects as controls (total 46 control eyes). Tears from 15 patients affected by allergic conjunctivitis (total 26 allergic conjunctivitis eyes) were also analyzed. Patients with systemic diseases (rheumatoid arthritis, liver diseases and cancer) that could alter the concentration of secreted HA were excluded from the study. The characteristics of the subjects are shown in Table I. The patients with unilateral acute adenovirus conjunctivitis (UAAC)
presented all of the following signs and symptoms: conjunctival hyperemia, ocular secretion, no corneal epithelium erosion, conjunctival follicular reaction, no use of any ocular medication and symptoms that had started for no more than 3 days prior. In the contralateral eye group, we included contralateral eyes from patients with UAAC that didn’t have any clinical signs of acute conjunctivitis. Diagnosis of allergic conjunctivitis was based on clinical history and evaluation of signs and symptoms. Patients included in the study were in an active inflammatory phase of the disease with active limbal infiltrates and were free of topical antihistamines and mast cell stabilizers for at least 3 days, topical corticosteroids for at least 7 days, and systemic antiallergic treatment for at least 2 week at the time of first presentations. The control group constituted of patients without any ocular or systemic diseases.

The ethics Committee of the Federal University of São Paulo approved this protocol and a written informed consent was obtained from each subject. The study protocol adhered to the tenets of The Declaration of Helsinki.

**SAMPLE COLLECTION**

For collecting the tears, Schirmer strips were placed in the temporal side of each eye under the eyelid, during 5 minutes, without any use of topical anesthetics. The strips were dried at room temperature and stored at ~20°C until analysis. The same procedure was performed for the three groups.

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<tr>
<th>TABLE I</th>
<th>Patients and donors characteristics.</th>
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<tr>
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<td>Patients UAAC</td>
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<tr>
<td>No.</td>
<td>15</td>
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<td>Age (mean ± SD)</td>
<td>32.6 ± 14.7</td>
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<td>Sex</td>
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<td>F</td>
<td>9</td>
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UAAC, unilateral acute adenovirus conjunctivitis; ALC, allergic conjunctivitis; F, female; M, male. Table shows standard errors for patients age.

**ADENOVIRAL CULTURE**

At the same time, conjunctival swabs were collected in the affected eye group in medium appropriate for adenoviral culture, using Hep-2 cells as described by de Paiva et al. 1992.

**TEAR SAMPLE PREPARATION**

Tear compounds were eluted from the Schirmer strips using 100 µL of distilled water, and HA and protein content analyses were performed.

**HA MEASUREMENT**

HA content in tear fluids was assayed by a non-competitive and non-isotopic fluoroassay (Martins et al. 2003). Eluted tear fluids and standard concentrations of HA (Sigma, St. Louis, MO) were added to 96 multiwell plates (FluoroNUNC Maxisorp-microtiterplates, Roskilde, Denmark) previously coated with HA-binding protein. The plates were then sequencially incubated with biotinylated HA-binding protein and europium-labeled streptavidin (Amershan, Piscataway, NJ). Afterwards, the europium remaining in the solid phase was released by an enhancement solution and the fluorescence was measured using a time-resolved fluorometer (Perkin–Elmer Life Sciences-Wallac Oy, Turkku, Finland). The data (counts/s) were processed automatically using the MultiCalc software program (Perkin-Elmer Life Sciences-Wallac Oy) and values are expressed as ng/mg protein.

**PROTEIN ANALYSIS**

Total tear protein concentration was determined using a colorimetric assay kit according to the manufacturer’s instructions (Protein Assay Kit from Bio-Rad, Hercules, CA). The protein profile was analyzed through sodium dodecysulfate polyacrylamide gel electrophoresis (SDS-PAGE) as previously described (Laemmli 1970). Briefly, 10 µg of protein from the tear samples were applied to a 3-20% linear gradient polyacrylamide gel under reducing conditions. After electrophoresis,
the gels were stained by comassie blue (Bio-Rad, Hercules, CA). Each protein band was quantified by densitometry using the software ImageJ Version 10.2 for Mac (U.S. National Institutes of Health, Bethesda, Maryland, USA). The results are expressed by arbitrary densitometric units (ADU).

STATISTICAL ANALYSIS

Data are expressed as mean ± standard error of the mean. Statistical analyses were performed using One-way ANOVA with Bonferroni’s Multiple Comparison test, using Graph Pad Prism 5.0 software for Mac. A 95% confidence interval and a 5% level of significance were adopted; therefore, results with P-value less than or equal to 0.05 were considered significant.

RESULTS

COMPARISON OF HYALURONIC ACID CONTENT IN TEAR FLUID FROM PATIENTS AFFECTED BY CONJUNCTIVITIS AND NORMAL DONORS

The HA content in tears from eyes with UAAC (n=15), from contralateral eyes (n=15), from eyes of normal donors (n=46) and from allergic conjunctivitis (n=26) was: 102.9 ± 19.9 ng/mg protein; 51.02 ± 8.5 ng/mg protein, 25.2 ± 2.3 ng/mg protein and 35.5 ± 4.5 ng/mg protein, respectively. The comparison of HA content among the three groups showed a significant difference (P<0.0001) (Fig. 1). UAAC tears presented higher HA concentrations when compared to either normal tears (P<0.0001), tears from contralateral eyes or tears from allergic conjunctivitis. HA content in tears from contralateral eyes did not differ (P>0.05) from tears from normal donors or from tears from patients affected by allergic conjunctivitis.

PROTEIN CONTENT AND QUALITATIVE ANALYSIS OF TEARS

The protein analysis showed distinct protein profiles when comparing tears from UAAC eyes to tears from contralateral eyes. Polypeptide bands were identified by matching their migrations to those of molecular weight standards. Tear protein patterns from normal donors have been exhaustively reported by previous investigators (Ballow et al. 1987, Lopez-Cisternas et al. 2006, Mann and Tighe 2007) and are represented by lactoferrin (79 kDa), slgA-heavy chain (66 kDa), slgA-light chain (27 kDa), lipocalin (TSPA, 18 kDa), and lysozyme (14 kDa) (Fig. 2). Figure 3 shows that eyes affected by UAAC presented higher amounts of slgA-heavy chain when compared to the contralateral eyes (CLE) or allergic conjunctivitis (ALC) (P=0.0368). On the other hand, UAAC presented lower quantities of lipocalin or TSPA (P=0.0394) and lactoferrin (P=0.0018) when compared to CLE. No differences were found in the protein content of slgA-light chain among UAAC, CLE and ALC. Finally, lysozyme is increased in tears from ALC compared to UAAC or CLE (P<0.0001).

DISCUSSION

We show that UAAC leads to a significant increase in tear HA content. The difference found between
HA tear content in UAAC was 2 times greater than that from contralateral eyes without clinical signs and 4 times higher than in the tears from normal donors indicating that HA could be a marker for subclinical inflammation. HA levels in tears of patients affected by allergic conjunctivitis showed no differences compared to tears from normal subjects.

Our results show that the levels of HA content in the tear fluid are related to the viral injury and conjunctival inflammation, suggesting that higher levels lead to a situation in which the viscoelastic properties of HA may protect corneal epithelium and help promote wound healing (Laurent and Fraser 1992, Miyauchi et al. 1996).

We may also consider that HA may be playing an important role in promoting inflammatory cell migration to the conjunctiva, possibly modulating inflammatory cytokine release (Mummert 2005).

UAAC, CLE and ALC tear protein profiles showed 79, 66, 27, 24, 18, and 14 kDa bands, which were a constant feature with different amounts of each protein (Fig. 2). The protein profile of tears is well established in normal eye (Kuizenga et al. 1991). In accordance to the data reported by other authors (Janssen and Van Bijsterveld 1981, Kuizenga et al. 1991), we propose that the proteins observed in the present study correspond to lactoferrin (79 kDa), slgA-heavy chain (66 kDa), slgA-light chain (27 kDa), lipocalin (TSPA, 18 kDa), and lysozyme (14 kDa), respectively.

A higher content of slgA was found in UAAC, when compared to CLE. Secretory IgA-enriched fluid presumably augments the effectiveness of the external barrier to microbial adherence and increases the efficiency by which pathogens are processed by the immune and inflammatory systems (Sack et al. 1992). No differences were found in slgA light chain content.

The decrease in lactoferrin and lipocalin levels in UAAC tears may occur evoked by major reflex tearing caused by inflammation (de Paiva et al. 1992). Furthermore, lactoferrin has been shown to

![Figure 2 - Protein profile in tear of allergic and adenoviral conjunctivitis. Electrophoretic profiles of human tear fluid on SDS-PAGE revealed by Coomassie blue staining. In A (contralateral eye), B (acute adenoviral conjunctivitis eye) and C (allergic conjunctivitis), the corresponding profiles of 4 different subjects are shown. Protein bands were identified by matching their migrations to those of molecular weight standards; lactoferrin (79 kDa), serum albumin and slgA-heavy chain (66 kDa), slgA-light chain (27 kDa), lipocalin (TSPA, 18 kDa), and lysozyme (14 kDa).]
possess antiviral activity against adenovirus. This possible down regulation may be an evolutionary mechanism used by adenovirus to survive in ocular tissue. There was no difference between UAAC and CLE regarding lysozyme secretion, but it was increased in ALC tears, since an allergen exposure can cause a marked increase in lysozyme secretion (Proud et al. 1998).

**CONCLUSIONS**

In this study we were able to identify an important increase in HA tear content in patients affected by UAAC, compared with normal donors or allergic conjunctivitis, using a rapid and non invasive method. Regarding protein analysis we could verify a distinct protein profile in tear fluid of UAAC patients. Taken together these results indicate that the increase in HA content seems to play a role in the tear fluid of UAAC patients. The quantification of HA in the tear fluid is a rapid, sensitive and specific test. This molecule might be a biomarker candidate for acute conjunctivitis.

**ACKNOWLEDGMENTS**

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**ABBREVIATIONS:**

UAAC, unilateral acute adenovirus conjunctivitis; CLE, contralateral non-affected eye; ALC, allergic conjunctivitis.
HYALURONIC ACID CONTENT IN CONJUNCTIVIS

conjunctivitis; HA, hyaluronic acid; GAG, glycosaminoglycan; SDS-PAGE, sodium dodecylsulfate polyacrylamide gel electrophoresis.

RESUMO

A conjuntivite causada por adenovírus é uma das maiores causas de infecção da conjuntiva no mundo. A conjuntivite provoca sintomas relativamente inespecíficos, como hiperemia e quemose. Mesmo depois de biomicroscopia, testes laboratoriais complexos, como cultura viral, são necessários para identificar o patógeno ou sua etiologia. Para contribuir para o melhor entendimento da fisiopatologia da conjuntivite causada por adenovírus, lágrimas de pacientes com conjuntivite aguda unilateral causada por adenovírus (UAAC), de doadores normais (controle) e de pacientes com conjuntivite alérgica foram analisadas. As amostras foram coletadas com tiras de Schirmer de doadores normais, pacientes com conjuntivite alérgica e pacientes com UAAC diagnosticados por sinais clínicos e testes positivos em culturas virais. Após a eluição das lágrimas, o HA foi quantificado utilizando um ensaio fluorométrico semelhante ao ELISA e o perfil da proteína foi determinado por SDS-PAGE. Um aumento profundo no conteúdo de HA em lágrima de pacientes com UAAC foi encontrado quando comparado com o controle ou a conjuntivite alérgica. Este aumento de HA em lágrimas de pacientes com UAAC não foi observado em lágrimas do olho contralateral sem sinais clínicos ou de pacientes com conjuntivite alérgica. Além disso observou-se um perfil distinto de proteínas nas lágrimas de pacientes com UAAC. A quantificação de HA no fluido lacrimal é um ensaio rápido, sensível e específico. Esta molécula pode ser um bom candidato a biomarcador para conjuntivite aguda.

Palavras-chave: conjuntivite, glicosaminoglicanos, ácido hialurônico, lágrima.

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