Acute hemorrhagic conjunctivitis: anti-coxsackievirus A24 variant secretory immunoglobulin A in acute and convalescent tear

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Purpose: The purpose of this paper is to present the clinical course of a laboratory-acquired case of acute hemorrhagic conjunctivitis (AHC) caused by coxsackievirus A24 variant (CA24v). Also, the anti-CA24v neutralizing activity and anti-CA24v immunoglobulin (Ig) G and secretory IgA (sIgA) in acute and convalescent tears and/or sera are presented.

Case: A 60-year-old male presented with acute-onset left eyelid edema, tearing, conjunctival erythema, pain, foreign body sensation, and subconjunctival hemorrhage 24 hours after suspected laboratory exposure. Bilateral conjunctivitis presented 24 hours later and resolved in 10 days.

Methods: Tear and blood samples were collected for virus isolation and neutralizing assays. CA24v-reactive IgG and sIgA in tear and/or serum samples were detected by immunofluorescent antibody analysis of ethanol-fixed virus-infected cells.

Results: Peak tear neutralization titers (1,000–1,500 U/mL) against the isolated virus occurred 1 day post-onset (po) of AHC. Tear neutralization titers became undetectable by the sixth day as serum neutralization titers became detectable on the ninth day po (60 U/mL), peaked by 21 days (3,000 U/mL), declined by 1 year to 200 U/mL, and remained at 30 U/mL 5 years po. Antibody to human IgG, IgA, and secretory component (sIgA) reacted with CA24v-infected cells treated with pooled acute tears collected 1–4 days po. Predominantly, sIgA was detected in CA24v-infected cells treated with tears collected 4 years and 5 years post-AHC, while convalescent serum contained predominantly anti-CA24v IgG.

Conclusion: AHC was confirmed by CA24v isolation, tear anti-CA24v neutralizing activity, and seroconversion. The detection of CA24v-reactive IgG, sIgA, and neutralizing activity in tears collected 1–4 days po of AHC supports plasma extravasation of IgG and suggests a defensive role for tear anti-CA24v sIgA. The results suggest that immunofluorescent antibody analysis of tears for persistent anti-CA24v sIgA may be useful in epidemiological monitoring of AHC.

Keywords: neutralization, immunofluorescence, eye infection, enterovirus, seroconversion

Introduction

Epidemics of acute hemorrhagic conjunctivitis (AHC) have occurred worldwide since the first cases were reported in Accra, Ghana, and Singapore in 1969–1970.1,2 Two picornaviruses, enterovirus 70 (EV70) and coxsackievirus A24 variant (CA24v), were isolated and shown to be the etiologic agents of AHC epidemics.2–4 AHC is highly contagious and is transmitted primarily by hand-to-eye-to-hand contact, but fomite and airborne spread is suspected. High attack rates in families and communities are common. High humidity, poor sanitation, overcrowded housing, and school/factory exposure facilitate spread during epidemics. AHC is endemic in several countries where sporadic cases occur annually and local outbreaks and epidemics...
The persistence of antibody to AHC viruses to post-infection is unknown.

We presented the clinical course of an AHC case through 6 days, demonstrating the acute increase in anti-CA24v neutralizing activity and secretory immunoglobulin (Ig) A (sIgA) in acute tears, and used immunofluorescence antibody analysis to show the persistence of CA24v-reactive sIgA in tears collected 4–5 years post-AHC. The detection of anti-CA24v sIgA in acute tears suggested local production and a defensive role against infection.

Case report

A 60-year-old male presented with foreign body sensation in the left eye and symptoms of conjunctivitis (Figure 1) 24 hours after possible laboratory exposure to AHC virus. His physical exam was within normal limits, and he reported no prior ocular infections, current systemic disease, or medication use. An external ocular examination of the left eye revealed lid edema, tearing, conjunctival erythema, dilated conjunctival blood vessels predominantly in the inferior temporal and nasal bulbar conjunctiva, and small blotches of subconjunctival hemorrhage in the inferior bulbar conjunctiva of the left eye (Figure 1A) that progressed to the infra-nasal bulbar conjunctiva on day 3 that resolved in 10 days without sequelae or treatment. No ocular infections were noted during the following 3 years. TEARS were noted, and the bilateral conjunctivitis resolved completely in 10 days without sequelae or treatment. No ocular infections were noted during the following 3 years. The severity of signs and symptoms peaked in the left eye on day 2 and in the right eye on day 3 post-onset (po).

The bilateral tearing, conjunctival erythema, and swelling increased through 24 hours, remained through 48 hours, and diminished by day 6 (Figure 1C and F). The right eye conjunctivitis mirrored that of the left eye, with onset of symptoms over 6 hours. The severity of signs and symptoms peaked in the left eye on day 2 and in the right eye on day 3 post-onset (po). The bilateral tearing, conjunctival erythema, and swelling increased through 24 hours, remained through 48 hours, and diminished by day 6 (Figure 1C and F). The right eye conjunctivitis mirrored that of the left eye, with onset of symptoms over 6 hours.

The factors promoting sporadic and recurrent AHC epidemics are unknown but likely include unrestricted travel,5,31 antigenic/genetic divergence of AHC viruses,32-34 subclinical infection,35,36 and the absence or waning of population immunity.3,37-39 An additional confounding factor is the low frequency of seroconversion (15%–60%) reported in many AHC epidemics, suggesting that the ocular infection may not provide protection.3,37-41 The weak systemic immune response to AHC is likely due to the highly localized nature of the infection to the eye (mucosal epithelium) and rapid viral reduction, but hyporesponsiveness or local antibody production exclusively has not been ruled out.42,43 Paradoxically, it has been reported that high pre-epidemic serum antibody titers provide sex-specific protection against EV70 AHC.44 Generally, subclinical infection is monitored between epidemics by detection of AHC virus and/or seropositive individuals, while the population’s relative susceptibility to an AHC epidemic is predicated upon the absence of, or decline in, serum antibody to the AHC viruses.1,32,33,37-41,44-46

Materials and methods

Tears and sera

Bilateral tear fluid samples (200 µL) for virus isolation, neutralization, and immunoreactive antibody studies were
collected one to two times daily in glass capillary pipettes 1–6 days po, and pooled bilateral tear samples (150 µL) were collected on 2 consecutive days at ~1 year, 4 years, and 5 years po as previously reported.\(^{23,25}\) Venous blood samples (1–3 mL) were collected at intervals between 2 days and 60 days, and at 4 months, 1 year, 4 years, and 5 years po using a 25-G infusion set with syringe and standard aseptic phlebotomy methods. Clarified tears and sera from clotted blood samples were collected after desktop centrifugation (1,000× g for 5 minutes) and were stored frozen (−10°C). Reference antisera obtained from the National Institutes of Health, Bethesda, MD, USA (NIH Research Reference Reagents) included antisera to CA24 (Joseph) (CA24/Africa/Joseph/1952; V027-501-563) and polioviruses type 1 (LSC; V001-511-560), type 2 (P-712; V002-511-560), and type 3 (Leon; V003-5110560). Antiserum to prototype CA24v (Singapore/EH24/1970) was provided by Dr M Yin-Murphy, Singapore University, Singapore. Antiserum to prototype EV70 (Japan/670/1971) was obtained from Reisaku Kono, National Institute of Health, Tokyo, Japan. This case report was reviewed and approved by the LSU Health Institutional Review Board, and the clinical investigations were conducted in accordance with the World Health Organization Declaration of Helsinki.

Cell culture

Human retinal pigmented epithelial (HRPE) cells (CRL-2502; ARPE-19) and African green monkey kidney (AGMK) cells were maintained as recommended by the provider; (American Type Culture Collection, Rockville, MD, USA). (It should be noted that cultured retinal pigmented epithelial cells express γFc receptor mRNA.\(^{47}\)) For experiments, trypsinized cells were suspended (5×10⁶ cells/mL) in Dulbecco’s minimal essential medium (DMEM; Sigma-Aldrich, St Louis, MO, USA) supplemented with 2% bovine calf serum (HyClone Laboratories, Logan, UT, USA) and antibiotics (100 U penicillin and 100 µg streptomycin/mL; Thermo Fisher Scientific, Waltham, MA, USA). Cell suspensions in DMEM were pipetted into six-well dish cultures (2 mL/well; Sarstedt AG & Co., Nümbrecht, Germany) for virus isolation and propagation, 96-well microtiter plate cultures (100 µL/well; Sarstedt AG & Co.) for neutralization assays, and Lab-Tek® eight-chamber glass slide culture (200 µL/well; Thermo Fisher Scientific, Rochester, NY, USA) for immunofluorescent antibody (IFA) analysis. The cultures were incubated for 24 hours at 37°C in a 5% CO₂-humidified atmosphere in a water-jacketed incubator (Forma-Scientific, Fredrick, MD, USA) until cell monolayers reached confluence.

Virus isolation and viruses

Tear fluid was applied directly to fresh medium over HRPE or AGMK cells and incubated at 37°C as previously
reported.\textsuperscript{23} Viral cytopathogenic effects in tear-inoculated cultures approached 100% after 24-hour to 48-hour incubation, and the culture media was harvested. Early-passage HRPE cell virus isolate sub-stocks were clarified (5,000x g for 10 minutes), aliquoted, and stored frozen (–80°C). Prototype (Singapore/SEC24/1970)\textsuperscript{2} and prime type CA24v Texas/M07/1977,\textsuperscript{23} prototype and prime EV70 types (Japan/ J670/1971 and Florida/KW97/1981, respectively),\textsuperscript{46} and/or poliovirus type 1 (Mahoney) were grown in HRPE cells and used in HRPE cell neutralization assays.

Neutralization assay

The neutralizing titers were determined from duplicate micro-neutralization assays in HRPE cells as previously described.\textsuperscript{23,25} Briefly, half-log\textsubscript{10} dilutions of tear and serum samples, as well as reference antisera, were reacted with 20–50 plaque-forming units (PFU) of Louisiana/LTV/2010, isolates of CA24v and EV70, and poliovirus serotypes.

The reciprocal of the mean endpoint dilution yielding 50% reduction in virus plaques was used as the neutralization titer (units per milliliter).

IFA assay

For all immunofluorescence experiments, HRPE and AGMK cells were grown on multiple Lab-Tek\textsuperscript{®} eight-chamber slides (Thermo Fischer Scientific), infected with 20–50 PFU of Louisiana/LTV/2010/culture and incubated at 37°C until plaques formed (usually 24 hours). The culture fluid was removed, and the cells were fixed with cold 70% ethanol (–10°C) for 30 minutes. The ethanol was removed, and the cells were washed three times with phosphate-buffered saline (PBS; pH 7.4). The cells were washed with PBS, and the nuclei were stained with 4',6-diamino-2-phenylindole (DAPI) as previously described.\textsuperscript{46} The DAPI stain was removed, and the cells were rinsed twice with PBS. Rabbit serum (1:100 dilution in PBS) was applied to block nonspecific binding and removed after 30-minute incubation at room temperature. Side-by-side duplicate slide culture wells on two slides were overlaid with 200 µL of a 1:30 dilution of pooled acute tears (collected 1–4 days po), 4-year or 5-year convalescent tears, or 1:100 dilution of an acute and convalescent serum in PBS. The tear and sera dilutions were removed after overnight incubation at 4°C, and the cells were washed three times with PBS. For dual IFA staining, 200 µL of PBS containing a 1:1,000 dilution of fluorescein isothiocyanate (FITC)-labeled goat anti-human IgG (Fc-chain specific) and tetramethylrhodamine isothiocyanate (TRITC)-conjugated goat anti-human serum IgA (Fc specific; Jackson ImmunoResearch Laboratories, Inc., West Grove, PA, USA) (diluted 1:1,000 in PBS) were added to duplicate slide culture chambers and incubated for 3 hours at room temperature in the dark. After removal of the secondary antibodies, the slides were rinsed six times with PBS, excess PBS was blotted off, and glass cover slips with mounting media were applied. To detect sIgA in acute and (4-year or 5-year) convalescent tears, a 1:1,000 dilution of goat antihuman secretory component (bound and free) (Sigma-Aldrich) in PBS was applied to tear-treated duplicate slide cultures for 6 hours at room temperature. Goat antibody to secretory component in tear-treated, virus-infected cells was detected by removing the tear/PBS, rinsing three times with fresh PBS, and applying a 1:1,000 dilution of TRITC-conjugated donkey anti-goat IgG (Jackson ImmunoResearch Laboratories, Inc.) in PBS and incubated at room temperature for 3 hours. After removal of fluorochrome-labeled secondary antibody, the slides were rinsed six times with PBS, excess PBS was blotted off, and glass cover slips with mounting media were applied. All IFA procedures and digital color images were generated as previously described.\textsuperscript{49} Red, green, and blue images were merged using Scanalytics IPLab 3.7 software.

Results

Virus isolation and serotyping

Cytopathogenicity consistent with enteroviral infection was detected in HRPE cell cultures 24 hours postinoculation with pooled bilateral tear samples (not shown). The isolated virus was designated Louisiana/LTV/2010 and was neutralized by rabbit antiserum raised against prototype CA24v (300 U/mL) but not by antisera to EV70, CA24 (Africa/Joseph/1952), or poliovirus types 1, 2, and 3 (not shown).

Tear neutralizing activity

Bilateral tear samples contained neutralizing activity against Louisiana/LTV/2010 (Figure 2). Neutralizing activity against the isolated virus was detected in left eye tears on the first day po but was not detected in tears from the fellow unaffected eye. Peak titers of tear neutralizing activity were detected in tears of the left (1,000 U/mL) and right (1,500 U/mL) eyes on days 1 and 2 po of AHC, respectively. Tear anti-CA24v neutralizing activity declined to near-undetectable levels bilaterally by day 6 and remained undetectable at 5 years. Concomitantly, neutralizing activity was detected in pooled 1-day to 4-day tears against prototype CA24v (300 U/mL) and poliovirus type 1 (20 U/mL) but not EV70 (<10 U/mL) (not graphed).
A greater than fourfold increase in the serum neutralizing antibody titer against the Louisiana/LTV/2010 isolate was detected between days 2 and 9 (10–60 U/mL). The serum neutralization antibody titer peaked at 3,000 U/mL on day 21 and declined from 2,000 U/mL at 4 months to 30 U/mL by 5 years (Figure 3). Serum neutralization titers against prototype CA24v Singapore/SEC24/1970 (10–600 U/mL) and prime type Texas/MO7/1977 (10–3,000 U/mL) paralleled those of the tear virus isolate. Anti-EV70 (Tokyo/J670/1971 and Florida/KW97/1981) neutralization was not detected in acute or convalescent serum (<10 U/mL), but anti-poliovirus type 1 (Mahoney) neutralizing antibody activity was detected in acute and convalescent sera (60–200 U/mL) (not graphed). Note that the sera neutralization titer was lower against the prototype CA24v Singapore/SEC24/1970 than the prime type CA24v Texas/MO7/1977 and the isolated virus, Louisiana/LTV/2010. Taken together, the results suggest that the isolated virus was antigenically similar to prototype and prime type CA24v.

**Detection of anti-CA24v IgA and IgG in tear and serum by IFA**

Anti-CA24v-reactive IgG and IgA were detected in ethanol-fixed, virus-infected HRPE and AGMK cells treated with pooled acute tears collected on 1–4 days and at 4 years or 5 years post-AHC as well as acute (11 days) and convalescent (5 years) sera (Figure 4). IgG and IgA in the pooled acute tears reacted with viral antigens in virus isolate-infected HRPE (Figure 4A) and AGMK cells (Figure 4B). Notably, some virus-infected cells were labeled with predominantly IgA or IgG. In some dual IgG- and IgA-labeled virus-infected cells, the IgA was generally localized to the cytoplasm (Figure 4A and B). Predominantly IgA, with little IgG, was detected in CA24v-infected cells treated with 4-year convalescent tears sample in HRPE cells (Figure 4C) or 5-year convalescent tear sample in AGMK cells (Figure 4D). Moreover, secretory component (indicative of sIgA) was detected in isolate-infected HRPE cells treated with pooled acute tears (Figure 4E) as well as 5-year convalescent tear (Figure 4F). In comparison, predominantly IgG labeling was detected in the isolate-infected cells treated with convalescent sera collected on 11 days (Figure 4G) and 5 years (Figure 4H). The results point to a predominance of anti-CA24v sIgA in convalescent tears and a predominance of anti-CA24v IgG in convalescent serum.

**Discussion**

The diagnosis of AHC was based upon history of recent AHC virus contact, short incubation period, acute onset of unilateral conjunctivitis with subconjunctival hemorrhage that spreads to the fellow eye within 48 hours, and complete resolution in 10 days.9 As in our case, the diagnosis of AHC is relatively simple for epidemic and laboratory-acquired infections. The etiological agent of the AHC was confirmed by isolation of a serotypic CA24v and demonstration of a fourfold increase in anti-CA24v neutralization titer between acute and convalescent serum samples. Further confirmation was CA24v-specific neutralizing activity in tears collected on 1–6 days.23,25,26
Figure 4: Immunoreactivity of tear and serum IgG (green) and IgA (red) from the AHC patient with CA24v-infected cells.

Notes: Arrows show similar differential staining patterns for tear IgG and IgA in CA24v-infected HRPE and AGMK cells. (A) Detection of immunoreactive human IgG and/or IgA in CA24v-infected HRPE cells treated with pooled acute tears (collected 1–4 days po). (B) Detection of immunoreactive human IgG and/or IgA in CA24v-infected AGMK cells treated with pooled acute tears. (C) Detection of immunoreactive human IgG and IgA in CA24v-infected HRPE cells treated with tears collected 4 years post-AHC. (D) Detection of immunoreactive human IgG and IgA in CA24v-infected AGMK cells treated with tears collected 5 years post-AHC. (E) Detection of antibody to secretory component (polymeric IgA) in CA24v-infected HRPE cells treated with pooled acute tears. (F) Detection of antibody to secretory component (polymeric IgA) in CA24v-infected HRPE cells treated with tears collected 5 years post-AHC. (G) Detection of CA24v immunoreactive human IgG and IgA in acute serum (collected 11 days po)-treated virus-infected HRPE cells. (H) Detection of CA24v immunoreactive human IgG and IgA in virus-infected HRPE cells treated with the patient’s convalescent serum (collected 5 years post-AHC). Bar = 50 µm.

Abbreviations: IgG, immunoglobulin G; IgA, immunoglobulin A; AHC, acute hemorrhagic conjunctivitis; CA24v, coxsackievirus A24 variant; HRPE, human retinal pigmented epithelial; po, post-onset; AGMK, African green monkey kidney.
The detection of anti-CA24v, but not anti-EV70, neutralizing activity with anti-CA24v IgG and sIgA in AHC tears collected through 6 days is consistent with the idea that inflammation, and innate and humoral immune responses act together to inhibit the virus infection.\textsuperscript{23–29} The detection of the neutralizing activity in early tears is consistent with Ig activity and the previously reported detection of serum protein and IgGs in 1-day to 3-day tears of epidemic AHC cases.\textsuperscript{30} The IFA detection of non-neutralizing anti-CA24v IgG in acute serum and anti-CA24v IgG in tear suggests serum IgG extravasation due to inflammation-induced increased vascular permeability. Extravasation of serum IgG into the tear is also supported by the detection of anti-polyovirus neutralizing activity (20 U/mL) in pooled acute tears. The concomitant IFA detection of IgA (and secretory component) in CA24v-infected cells treated with pooled acute tears in the face of very low serum levels of anti-CA24v IgA suggests local ocular production of IgA.\textsuperscript{51,52} Moreover, the anti-CA24v sIgA was detectable by IFA in tears at 5 years post-AHC. Interestingly, the detection of anti-CA24v sIgA with neutralizing activity in acute, but not in convalescent, tears suggests that the local ocular response shares similarities with the systemic responses reported for rubella, measles, and varicella-zoster viruses, in that virus-specific IgA first appears as neutralizing IgA and later becomes, or is replaced by, non-neutralizing IgA.\textsuperscript{53} The detection of anti-CA24v sIgA and neutralizing activity in acute tears suggests that sIgA may neutralize extracellular virus and/or block virus entry. This idea is consistent with the emerging concept that polymeric IgA antibody may be taken into the cell during virus infection where it triggers intracellular virus neutralization and innate immune signaling.\textsuperscript{51,54,55}

Alternatively, the possibility that cross-reactive neutralizing antibody\textsuperscript{30} or co-binding of IgG and IgA to CA24v may account for the early neutralizing activity in acute tear cannot be ruled out. Additional investigations are needed to determine if the early tear neutralizing activity is due to sIgA neutralization of CA24v and whether convalescent tear anti-CA24v sIgA protects against infection.

The levels and patterns of tears IgG and IgA labeling of CA24v-infected cells differed from that of serum. There is a possibility that polyclonal human IgG and IgA react with different viral epitopes/antigens. (Note: tear and serum IgG and IgA bound to virus antigens were probed simultaneously.) That is, virus-infected cells treated with pooled acute tear exhibited IgG and/or IgA labeling, with some dual labeled cells exhibiting dense cytoplasmic IgA with peripheral IgG (Figure 4A and B). In contrast, virus-infected cells treated with convalescent (4 years or 5 years) tears exhibited predominantly cytoplasmic sIgA labeling. The strong IgA and weak IgG signals within CA24v-infected HRPE cells treated with convalescent tear collected 4 years po may be due to a higher concentration of IgA in normal tears,\textsuperscript{37} while the lower IgA signal in 5-year convalescent tear-treated cells could represent a reduction in the anti-CA24v IgA. Moreover, the anti-CA24v sIgA signal in virus-infected cells treated with convalescent tear supports the persistence of CA24v-specific IgA years after infection.\textsuperscript{38} Concomitantly, the anti-CA24v IgG IFA signal and neutralizing activity in convalescent serum at 5 years post-AHC supports the persistence of anti-CA24v IgG.

**Conclusion**

In summary, the acute onset, signs and symptoms, and bilateral course in a previously seronegative laboratory employee were consistent with AHC epidemic cases\textsuperscript{16,30} and previously reported laboratory-acquired AHC virus infections.\textsuperscript{23,59} The results of the tear and serum analysis support the rapid appearance of anti-CA24v IgG, sIgA, and neutralizing activity in tears prior to the serum antibody response. The anti-CA24v sIgA in pooled acute tears suggests that it may play a role in limiting the CA24v infection. The persistence of anti-CA24v sIgA in tear up to 5 years suggests that it may have value in monitoring population immunity to CA24v AHC. Tear collection is tolerated well and provides useful diagnostic, epidemiological, and immunological information. Finally, the occurrence of this inadvertent laboratory infection underscores the necessity for strict conformity to standard aseptic techniques and pre- and post-use decontamination of shared work areas and equipment to eliminate AHC virus infection of laboratory personnel.

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