Impression cytology and in vivo confocal microscopy in corneas with total limbal stem cell deficiency

Citologia de impressão e microscopia confocal in vivo em córneas com deficiência total das células-tronco do limbo

Aline Lütz de Araújo1, José Reinaldo da Silva Ricardo1, Vivian Naomi Sakai1, Jeison Nadai de Barros1, José Álvaro Pereira Gomes1

ABSTRACT

Purposes: To describe corneal changes seen on in vivo confocal microscopy in patients with total limbal stem cell deficiency and to correlate them with cytological findings.

Methods: A prospective case series including 13 eyes (8 patients) with total limbal deficiency was carried out. Stem cell deficiency was diagnosed clinically and by corneal impression cytology. Confocal images of the central cornea were taken with the Heidelberg Retina Tomograph II, Rostock Corneal Module (Heidelberg Engineering, Heidelberg, Germany).

Results: Impression cytology of the cornea revealed conjunctival epithelial cells and goblet cells in all cases. In vivo confocal microscopy showed disruption of normal layers of the corneal epithelium in all eyes. Confocal images showed cells with characteristics of conjunctival epithelium at the cornea in 76.9% of the total. These findings on confocal microscopy were compatible to limbal stem cell deficiency. Additionally, goblet cells, squamous metaplasia, inflammatory cells and dendritic cells were observed. The sub-basal nerve plexus was not identified in any of the corneas. Corneal neovessels were observed at the epithelium and stroma. All cases showed diffuse hyper-reflective images of the stroma corresponding to opacity of the tissue.

Conclusions: Limbal stem cell deficiency had been confirmed by impression cytology in all cases, and 76.9% of the cases could also be diagnosed by in vivo confocal microscopy through the conjunctival epithelial cell visualization on the corneal surface. Frequent confocal microscopy findings were abnormal cells at the cornea (conjunctival epithelial, goblet and inflammatory cells), corneal neovessels and diffuse hyper-reflection of the stroma.

Keywords: Cornea/cytology; Corneal diseases; Conjunctiva/cytology; Limbus corneae; Stem cells; Microscopy, Confocal

INTRODUCTION

Limbal stem cell deficiency (LSCD) is a clinical condition that occurs in a variety of ocular diseases, including ocular chemical injuries, Stevens Johnson syndrome and cicatricial pemphigoid. Typical LSCD symptoms are decreased vision, photophobia, tearing, burning and recurrent pain episodes. The clinical picture is a conjunctivalized corneal surface with neovessels and recurrent or persistent epithelial defects. Tests for LSCD should demonstrate the presence and the extension of the deficiency. Other tests routinely performed target associated conditions like dry eye and neuroanatomical changes of the corneal surface. These tests include biomicroscopy, ocular surface staining with fluorescein, Schirmer tests, tearfilm breakup time, and corneal esthesiometry. Impression cytology is used to demonstrate goblet cells on the corneal surface and confirm the diagnosis of LSCD. However, it is important to consider the possibility of a false-negative diagnosis of LSCD when the presence of goblet cells is the only criterion considered, since a lack of these cells can happen on the sample by chance. More recently, in vivo confocal microscopy (IVCM) made possible to evaluate in vivo all corneal layers, from the epithelium to endothelium and its use to diagnose LSCD has already been attempted. IVCM shows corneal structures at different depths through the light reflected from the tissue. As

Submitted for publication: January 16, 2013
Accepted for publication: June 11, 2013

Study carried out at Department of Ophthalmology, Universidade Federal de São Paulo, São Paulo (SP), Brasil.

1 Physician, Universidade Federal de São Paulo, São Paulo (SP), Brasil.

Funding: No specific financial support was available for this study.

Disclosure of potential conflicts of interest: A.L.de Araújo, None; J.R.S. Ricardo, None; V.N.Sakai, None; J.N.de Barros, None; J.A.P.Gomes, None.

Corresponding address: Aline Lütz de Araújo, Rua Professor Cristiano Fischer, 34 - Porto Alegre (RS) - 90410-000 - Brazil - E-mail: alinelutz.a@gmail.com

Ethical approval: Project number 0287/10 approved by the institutional Ethics Committee in Research.
Impression cytology and in vivo confocal microscopy in corneas with total limbal stem cell deficiency

Visonest Heidelberg, Germany. A drop of anesthetic (proxymetacaine 0.5%, Tomograph II, Rostock Corneal Module (Heidelberg Engineering, analyzed under light microscopy. (Visonest to the conjunctiva with forceps after topical proxymetacaine 0.5% were cut in triangle shapes for orientation purposes and then applied pore size of 0.45 micron (Millipore Corporation, Bedford, USA). Filters in 4 quadrants of the corneal surface). after phototherapeutic keratotomy (20).

METHODS

We conducted a cross-sectional study with patients with total LSCD from the Cornea and External Disease Section of the Department of Ophthalmology, Universidade Federal de São Paulo, in Brazil. Ethical approval was obtained from the Institutional Medical Ethics Committee and National Ethics Committee in Research and followed the Tenets of the Declaration of Helsinki. Informed consent was obtained from the subjects after explanation of the nature and possible consequences of the study. The inclusion criteria were diagnosis of total LSCD based on the presence of conjunctival epithelial ingrowth onto the cornea in 4 quadrants (clinical signs observed are loss of corneal epithelial transparency, superficial corneal neovascularization, and epithelial irregularity or recurrent epithelial breakdown) (12) and on impression cytology (presence of goblet cells in 4 quadrants of the corneal surface).

For impression cytology, we used cellulose acetate filters with a pore size of 0.45 micron (Millipore Corporation, Bedford, USA). Filters were cut in triangle shapes for orientation purposes and then applied to the conjunctiva with forceps after topical proxymetacaine 0.5% (Visoneset®, Allergan, São Paulo, Brazil) instillation. Filters were pressed for 5 seconds onto the 4 quadrants of the cornea and then the specimens were fixed and stained with polyglandular autoimmune syndrome (PAS), hematoxylin and modified Papanicolau’s staining, according to previously described technique (21). Each specimen was analyzed under light microscopy.

IVCM of the cornea was performed using the Heidelberg Retina Tomograph II, Rostock Corneal Module (Heidelberg Engineering, Heidelberg, Germany). A drop of anesthetic (proxymetacaine 0.5%, Visoneset®, Allergan, São Paulo, Brazil) was instilled in the eye being examined. The object lens of the microscope was covered with gel (poliacrylic acid 0.2%, Vidisic Gel®, Bausch & Lomb, Berlin, Germany) to contact the cornea. Images were acquired from the central cornea at different depths, from the superficial epithelium to the endothelium. A digital camera placed on the side of the apparatus, giving a lateral image of the patient’s eye, controls eye position during examination.

We analyzed the following parameters: corneal epithelium characteristics (presence of normal cells, metaplastic cells, conjunctival epithelium cells, goblet cells, inflammatory cells, dendritic cells, neovessels); sub-basal nerve plexus with nerve density; stromal cells and neovessels; and endothelial cells.

Statistical analyses were performed using the Statistical Package for the Social Sciences 13.0 (SPSS Inc., Chicago, IL). Wilcoxon test was used to compare cytology and confocal microscopy.

RESULTS

Thirteen eyes of 8 patients (4 males) were evaluated. Mean age was 48.12 ± 17.8 years. The causes of LSCD (and the number or patients and eyes, respectively) were: ocular burn (1 patient, 2 eyes), Staphylococcia (1,2), aniridia (1,2), Stevens Johnson syndrome (1,1), polyglandular autoimmune syndrome (1,1), multiple surgeries (1,1), e idiopathic (2,4).

IMPRESSION CYTOLGY

According to inclusion criteria in the study, all cases had goblet cell on cytology samples. Also, all samples showed the presence of conjunctival epithelial cells on the corneal surface, 76.9% (10 cases out of 13) showed squamous metaplasia, and 69.2% (9/10) showed inflammatory cells in cytological analysis (Figures 1 and 2).

IN VIVO CONFOCAL MICROSCOPY

IVCM demonstrated that none of the cases had normal corneal epithelial structure (Figures 1 e 2). The basal layer of the corneal epithelium with its cells with bright borders arranged in a regular mosaic

Figure 1. Idiopathic limbal stem cell deficiency, A) biomicroscopy; B) impression cytology with inflammatory cells and conjunctival epithelial cells; C) in vivo confocal microscopy with corneal and conjunctival epithelia; and D) in vivo confocal microscopy with dendritic cells.

Figure 2. Limbal stem cell deficiency due to multiple surgeries. A) biomicroscopy; B) impression cytology with inflammatory cells, conjunctival epithelial cells bundle, and goblet cells; C) in vivo confocal microscopy with inflammatory and conjunctival epithelium cells; and D) in vivo confocal microscopy with superficial neovessels.
was seen in 23.1% (3 cases out of 13) of the cases by IVCM. No cell could be identified by the IVCM exam, aside from inflammatory cells, at any depth of the cornea in 23.1% (3/13). Once IVCM depends on light reflection from the tissue being examined, light reflection may become too scattered when cornea opacity is severe. In this situation, the images are rather blurred that no cell can be identified. Although these cases had been diagnosed with LSCD by impression cytology, the images obtained from IVCM were not helpful since no cells could be visualized.

IVCM identified cells suggestive of conjunctival epithelial cells on the cornea in 76.9% (10/13) and goblet cells (typically small, round and hypo-reflective cells) in 69.2% (9/13). Epithelial squamous metaplasia (seen as larger epithelial cell with diminished nucleus) was present in 15.4% (2/13). Round hyper-reflective particles suggestive of inflammatory cells were seen in all cases at different epithelial layers. Dendritic hyper-reflective bodies that characterize Langerhans cells were detected in 30.8% (4/13), all of which had idiopathic LSCD. The sub-basal nerve plexus was not identified in any of the corneas. Neovessels were seen at the epithelium in 84.6% (11/13) and at the stroma in 53.8% (7/13). Inflammatory cells at the stroma were seen in 7.7% (1/13), in all other cases images were diffusely blurred at the stromal level, probably due to stromal opacity. Endothelial cells were not visualized in any case.

**Impression cytology versus in vivo confocal microscopy**

Cytological and IVCM findings for each patient are presented in table 1. In 76.9% of the total (10 cases out of 13) the 2 exams agreed on LSCD diagnosis, since both detected conjunctival epithelial cells on the corneal surface. The number of eyes diagnosed as LSCD based on the presence of conjunctival cells on impression cytology or IVCM was not statistically different (p=0.08). There was a difference in the ability of the 2 exams in detecting goblet cells (p=0.04), squamous metaplasia (p<0.01), and inflammatory cells (p=0.04). For the 2 former findings, impression cytology showed better performance, and for the latter, IVCM performed better.

**DISCUSSION**

A number of alterations were found by corneal IVCM in patients with LSCD. We highlight the finding that corneal layers were not seen in full in most cases. Normal corneal epithelium has superficial, intermediate and basal layers. Since in LSCD this normal composition is not seen and conjunctival epithelial cells are found on the cornea, it is rational to conclude that conjunctival epithelium is replacing corneal epithelium. In some cases, IVCM showed both corneal and conjunctival epithelium at the same image. Nevertheless, even when there was some corneal epithelium on total LSCD, its structure was not normal. IVCM could not detect the sub-basal nerve plexus in our group of patients. Vera et al., examined 15 eyes with LSCD due to Stevens Johnson syndrome and were unable to identify the subbasal nerves on IVCM. It was not clear if the plexus was absent or was just undetectable because of the major modifications associated with the growth of neovessels in the basal section.

Impression cytology is the sampling technique of choice for ex vivo evaluation of superficial corneal and conjunctival cells, since it is minimally invasive and can be available for routine clinical use. The surface epithelium is the target tissue of interest rather than the basal epithelium or basement membrane. LSCD is confirmed by impression cytology through goblet cells identification on the cornea, although their absence on the smear may happen by chance and thus does not exclude the diagnosis. Immunohistochemistry may be used on impression cytology specimens to increase the accuracy. While healthy corneas express cytokinin keratin 12 (K12), corneas from patients with LSCD are characterized by the presence of mucin 1 (MUC1) and keratin 19 (K19).

In our study, IVCM findings in corneas with total LSCD were compared to impression cytology findings in order to evaluate IVCM as a tool for LSCD diagnosis. In 10 out of 13 eyes, IVCM was able to detect the presence of cells that were morphologically identical to conjunctival cells on the cornea, thus we found an agreement between the 2 exams on the diagnosis. For the remaining 3 eyes, IVCM could not detect any type of cell due to the haziness of the images (except for inflammatory

| Table 1. Impression cytology and in vivo confocal microscopy findings in each patient |
|---------------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Patient (OD/OS) | Diagnosis | Goblet cells | Conjunctival cells | Squamous metaplasia | Inflammatory cells |
| OD/OS | IC | IVMC | IC | IVMC | IC | IVMC | IC | IVMC |
| 1 (OD) | Chemical injury | + | + | + | + | + | + |
| 1 (OS) | Chemical injury | + | + | + | + | + | + |
| 2 (OD) | Idiopathic | + | + | + | + | + | + |
| 2 (OS) | Idiopathic | + | + | + | + | + | + |
| 3 (OD) | Idiopathic | + | + | + | + | + | + |
| 3 (OS) | Idiopathic | + | + | + | + | + | + |
| 4 (OD) | Staphylococcia | + | + | + | + | + | + |
| 4 (OS) | Staphylococcia | + | + | + | + | + | + |
| 5 (OD) | Aniridia | + | + | + | + | + | + |
| 5 (OS) | Aniridia | + | + | + | + | + | + |
| 6 (OD) | PAS | + | + | + | + | + | + |
| 7 (OD) | SJS | + | + | + | + | + | + |
| 8 (OD) | Multiple surgeries | + | + | + | + | + | + |

OD= right eye; OS= left eye; PAS= polyglandular autoimmune syndrome; SJS= Stevens Johnson syndrome; IC= impression cytology; IVCM= in vivo confocal microscopy.

The symbol + indicates the feature's presence.
cells, that are highly hyper-reflective). Therefore, IVCM did not refute LSCD diagnosis in those cases; it actually was not able to analyze the cells. Light scatter from extremely opaque corneas may limit the quality of the images when corneal opacity is too dense, although moderate or moderate to severe opacity is not a restriction for the exam. Our data showed an agreement between IVCM and impression cytology on LSCD diagnosis in all cases but those 3 in which IVCM images did not show cellularity, supporting that IVCM may be considered a valuable tool for the noninvasive in vivo diagnosis of LSCD.

Some findings were seen on IVCM and not on cytology, including hyper-reflective particles suggestive of inflammatory cells hyper-reflective bodies suggestive of dendritic (Langerhans) cells. The latter are deeply situated at the epithelium and basal membrane so they are not harvested by the impression sample. If located on central cornea, dendritic cells are indicative of inflammation and are associated with loss of “immune privilege” of the anterior segment, exacerbation of herpetic and other infectious keratitis and amplification of transplant immunity. In vivo confocal microscopy thus offer additional information, because impression cytology does not reach deeply located cells, such as inflammatory and Langerhans, that are indicative of corneal inflammation. Also, stromal neovessels and inflammatory cell on the stroma are seen on IVCM but not on impression cytology. All these information are helpful in the clinical context. Conversely, impression cytology as an ex vivo examination allows the use of staining to better evaluate epithelial cell changes and immunohistochemistry techniques to identify corneal and conjunctival cytokeratins. Therein, IVCM and impression cytology may complement one another. In summary, pros of IVCM are the visualization of deeper tissues, including cells and nerves of all corneal layers, and the possibility of repeated exams over time; at the opposite, as an in vivo exam, it does not allow staining of cells. Another con we found was that IVCM images weren’t adequate in cases of severe corneal opacity. Pro of impression cytology is the possibility of cell staining, including immunostaining, and con is that only the surface epithelium is harvested. Impression cytology may also be more available than clinoscopes because 5 eyes with suspect corneal conjunctivalization and found that IVCM and immunofluorescence CK12 staining of the cells from impression cytology were concordant on the diagnosis in 87% of the cases. Other study enrolled 58 eyes with ocular chemical injury; IVCM found goblet cells in 9 of them while impression cytology had the same result in 8 eyes. A positive correlation between goblet cell densities in the 2 exams was demonstrated. Adequate correlation between the IVCM and impression cytology was also demonstrated in dry eye and rosacea-related epitheliopathy. Common findings for both exams in dry eye are squamous metaplasia, inflammatory cell infiltration, goblet cell depletion, as well as a nuclear snake-like chromatin pattern. In rosacea associated with corneal epitheliopathy, goblet cells together with conjunctival epithelial cells within the corneal epithelium layer were found.

CONCLUSION

In our study, frequent IVCM findings in total LSCD were abnormal cells at the corneal epithelium (conjunctival epithelial, goblet and inflammatory cells), corneal neovessels and diffuse hyper-reflection of the stroma. Normal corneal epithelium comprising superficial, intermediate and basal layers was not seen in any case. LSCD had been confirmed by impression cytology in all cases, and 76.9% of the cases could also be diagnosed by IVCM through the conjunctival epithelial cell visualization on the corneal surface. We also find a limitation of IVCM in total LSCD when corneal opacity is severe. In those cases, confocal images were rather blurred that no cell can be identified. As a noninvasive technique, IVCM may be considered a valuable tool for LSCD diagnosis, and to follow up the condition and its response to treatment. In our study, patients had total LSCD. Studies with partial LSCD are needed to demonstrate the usefulness of IVCM in less severe cases.

ACKNOWLEDGEMENTS

We thank Dr. Charles McGhee, MD and Dr. Dipika Patel, MD from the Department of Ophthalmology at Auckland University New Zealand for helping us analyze confocal images.

REFERENCES