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Publication Date

2017-07-05

DOI

10.3791/55591

Peer reviewed

Video Article A Non-invasive Way to Isolate and Phenotype Cells from the Conjunctiva

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URL: https://www.jove.com/video/55591 DOI: doi:10.3791/55591

Keywords: Immunology, Issue 125, Impression cytology, dry eye disease, impression cytology device, effector memory T cells, central memory T cells, terminally differentiated effector memory T cells, HLA-DR, ocular surface impression cytology, flow cytometry

Date Published: 7/5/2017

Citation: Bose, T., Hou, A., Lee, R., Tong, L., Chandy, K.G. A Non-invasive Way to Isolate and Phenotype Cells from the Conjunctiva. J. Vis. Exp. (125), e55591, doi:10.3791/55591 (2017).

Abstract

Traditionally, ocular surface cytology is studied with techniques such as spatula technology and brush technology. The problem with these techniques is that they may induce traumatic lesions on the surface of the eye, which can progress to scarring, eyelid deformity, limbal stem cell deficiency and in some cases, cause great discomfort to the subject. To avoid these clinical problems, impression cytology (IC) was developed to diagnose dry eye disease and later neoplasia, atopic disease, vernal keratoconjunctivitis and keratoconjunctivitis sicca. Typically, clinicians manually cut filter papers into required shapes and apply these to the ocular surface. Here, we describe how to perform IC using a commercially available medical device. This technique is explained here followed by immunophenotyping by flow cytometry. This technique requires less manual handling and causes less injury to the ocular surface.

Video Link

The video component of this article can be found at https://www.jove.com/video/55591/

Introduction

Impression cytology (IC) was first performed in 1977 by *Thatcher et al*¹. They used a plastic impression disc to collect conjunctival cells from patients instead of other techniques available at that time such as scraping, swabbing or pipetting¹. The current technique of IC uses an absorbent filter paper² to imprint the bulbar and palpebral conjunctiva and collect the most superficial layer of conjunctival cells. These cells, having reached their final stage of differentiation, are continuously shed into tears³. Three major populations of cells are found in IC specimens: epithelial cells⁴, goblet cells^{3,5}, and mucosal-associated lymphoid tissue *viz* epithelium-associated effector T cells or dendritic cells⁶. Ocular surface cells in IC samples can be analyzed by microscopy, immune-blotting and reverse-transcriptase polymerase chain reaction (RT-PCR)⁷. Flow cytometry has been recently used to analyze immune cells collected by scraping the IC membrane⁸. Interestingly, IC^{6,9} has been used to evaluate many ocular surface diseases including keratoconjunctivitis sicca, vitamin A deficiency, cicatricial pemphigoid, atopic disease, superior limbic keratoconjunctivitis, vernal keratoconjunctivitis, and epithelial squamous metaplasia. IC has also been used to evaluate the impact of wearing contact lenses, detecting ocular surface microbes, and testing therapeutic efficacy and tolerance of therapeutic interventions in longitudinal studies^{10,11,12}.

The medical device (EyePrim) is supported by a type of polyethersulfone (PES) 0.2-µm membrane, which has been previously validated for the technique of ocular impression cytology with flow cytometry (OSIC-flow) and opens up opportunities to use longitudinal sampling to monitor disease progression and response to treatment (*e.g.*, the detailed analysis of intraepithelial leukocytes defined as putative disease markers for progressive conjunctival fibrosis in mucous membrane pemphigoid)¹³. Early researchers used autoclaved PES filters that required manual impression. As a result, the yield was variable and user dependent. The advantage of this medical device is ease of use, standardized pressure (Pa or N/m²), and enables repeatability, reproducibility, and consistent cellular recovery. This technique is useful in an out-patient clinic because it is non-surgical, easy-to-perform and rapid. This is a Class I (sterile) medical device according to the directive 93/42/CEE, CE 0499 (SNCH). It only requires topical anesthesia during the procedure, which ensures maintenance of the integrity of the ocular surface. Following IC, cells can be processed immediately for flow cytometry. Moreover, it is possible for non-ophthalmology technicians and nurses to be trained to sample the ocular surface.

Despite the improvement of IC over other techniques, several challenges remain. For example, there may be variation due to the area of the sampling and regional differences in the bulbar conjunctiva depending on the position of the IC. Another source of variation is due to the application of different amounts of pressure during IC. Other methodological issues involve standardization of cell processing: these involve duration and method of fixation, and the conditions of possible storage, which may impact stability of the sampled material.

The overall goal of this technique is to develop a method of isolation of the ocular impression samples that is easy to use, non-invasive and can be applied to the immunological characterization of the clinical samples.

Protocol

All ocular samples used in this study were collected from the Dry Eye Clinic at the Singapore National Eye Centre approved by SingHealth Centralized Institutional Review Board and the Nanyang Technological University Institutional Review Board, Singapore.

NOTE: Subjects underwent clinical tests to assess the extent of inflammation and the severity of tear dysfunction before IC was performed. Clinical tests included non-invasive tear break-up time (NI-TBUT)¹⁴ and conjunctival redness (hyperemia)^{15,16} assessed with a diagnostic instrument, Schirmer's test¹⁷, Standard Patient Evaluation of Eye Dryness (SPEED) questionnaire¹⁸, and corneal staining¹⁹.

1. Collection of Ocular Samples by IC

- 1. After determination of the clinical status, anesthetize the participant's eyes by applying 1 2 drops of topical anesthesia, proparacaine hydrochloride, to the superior bulbar conjunctiva and inferior fornix. Wait for 4 5 min for the stinging sensation of the anesthetic to wear off.
- Collect samples from both nasal and temporal bulbar conjunctiva with two impression cytology devices. NOTE: One device is used for the collection of two impression samples. Here, two devices were used for the collection of four impression membranes.
 - Position the impression cytology device tangentially on the conjunctiva. Press the push-button gently and hold for 2 3 s. NOTE: The device comes with the membrane. Pressure is released automatically after pushing the button. It takes only few seconds to release the pressure and remove the device.
- 3. Release the membrane into a microcentrifuge tube containing 1 mL of culture media (Roswell Park Memorial Institute medium (RPMI) + 10% fetal bovine serum (FBS) + 1% Penicillin-Streptomycin (Pen Strep)) by a scalpel.
- Isolate cells from the membrane of the device by continuous scraping for around 1 min with a 10 μL pipette tip. NOTE: The scraping is completed when the surface of the membrane becomes uneven. The cell numbers from each membrane is variable (100 - 500 cells/membrane). Process the samples within 2 - 3 h of collection.

2. Immunophenotyping Cells by Flow Cytometry

- Centrifuge the cells at 400 x g for 5 min at room temperature to remove the media. Resuspend the cell pellet with 50 μL of flow cytometry staining buffer (phosphate buffered saline (PBS) + 0.05% bovine serum albumin (BSA)).
- 2. Stain cells with a panel of antibodies (*e.g.*, CD3 brilliant violet (BV) 510 (UCHT1), CD4 allophycocyanine-H7 (APC-H7) (SK3), CCR7 phycoerythrin (PE)-A, CD45RO PE-cyanin 7 (PE-Cy7)-A, and live/dead cell mark 7-ADD) at room temperature for 20 30 min in the dark. Use antibodies directly from the stock according to the manufacturer's protocol. NOTE: For the antibody concentration, use 2.5 µL of CD3-BV510, 1.25 µL of CD4-APC-H7, CD45RO-PE-Cy7, 7-AAD and 10 µL of CCR7-PE. Use peripheral blood mononuclear cells (PBMCs) to titrate the different concentrations of antibodies. For the titration, take the antibody concentration mentioned in the manufacturer's protocol and use half and one-fourth of the recommended concentration. Use the minimum concentration of antibodies to avoid spill-over to other fluorochrome channels. Clear signals were observed with the mentioned concentrations of antibodies. The compensation was performed by testing PBMCs initially according to the protocol mentioned in Williams *et al.*²⁰ For testing, PBMCs were incubated with the same set of antibodies used here and compensated by comparing single and multiple antibody stains.
- 3. After the incubation period, add 1 mL of staining buffer to the tube, mix well, and centrifuge at 450 x g for 5 min at room temperature. Resuspend cells in 200 µL of staining buffer.
- 4. Run samples on a flow cytometry machine.
 - 1. Before acquiring the data, calibrate the flow cytometer channel voltages with cytometer setup and tracking (CS&T) beads to normalize the data acquisition on different days as per manufacturer's instructions.
 - 1. Open the flow cytometry data collection software, click "Set up & QC" button, choose the correct CS&T bead lot number, load the beads, and then click the "Start" button.
 - Resuspend cells by gently tapping the tube before loading samples to the machine. Open the data collection software and click "Preview". When threshold rate is stable, click "Acquire". Monitor the sample level and click "Stop" when samples run out. NOTE: Acquire 10,000 events per sample.
 - 3. After acquiring 10,000 events, generate a SSC-A and FSC-A dot plot in the worksheet.
 - Click the "Polygon Gate" button and draw a gate (this is P1) over all events with a FSC-A value > 5 x 10⁴ to exclude debris. Click "Create dot plot" button to generate a new dot plot, right click the dot blot, choose "Properties" to open the "Plot Editor" window, and then choose "P1". Click the FSC-A on x-axis of P1 dot plot, change it to CD3-BV510. Draw a "Polygon Gate" to choose the CD3⁺ population and name it "P2".
 - Click "Create dot plot" button to generate a new dot plot, right click the dot blot, choose "Properties" to open the "Plot Editor" window. Click the FSC-A on x-axis of P2 dot plot and change it to 7-AAD. Draw a "Polygon Gate" for 7-AAD⁻ population and choose "P3". P3 here is CD3⁺ live cells.
 - 3. Click "Create dot plot" button to generate a new dot plot, right click the dot blot, choose "Properties" to open the "Plot Editor" window. Click the CD3-BV510 on x-axis and CD4-APCH7 on y-axis of the P3 dot plot. Draw Rectangle Gates on the upper and lower sides of the quadrant and choose "P4" and "P5", respectively. P4 is live CD3⁺CD4⁺ and P5 is live CD3⁺CD4⁻ T cells.
 - 4. Click "Create dot plot" button to generate a new dot plot, right click the dot blot, choose "Properties" to open the "Plot Editor" window. Click the CD45RO PE-Cy7 on the x-axis and CCR7-PE on the y-axis of both the P4 and P5 plots. Draw "Quad Gate" on this dot plot.

NOTE: Upper left hand side, upper right hand side, lower right hand side and lower left hand side quadrants here are naïve, central memory (T_{CM}), effector memory (T_{EM}), terminally differentiated effector memory (T_{EMRA}) T cells, respectively.

Representative Results

IC by this clinical device allowed us to isolate ocular surface immune cells while keeping the ocular surface intact. **Figure 1** describes how the IC was performed. The different sites of the eye from where the samples were collected are marked. **Figure 2** shows representative result of flow cytometry collected from 10 healthy controls. For the gating, use SSC and 7-AAD to distinguish between live and dead cells. 7-AAD⁻ cells are identified as live cells. These live cells are further characterized by CD3⁺ and CD4⁺ markers. Furthermore, the CD4⁺ and CD4⁻ populations were each further characterized as Naïve, T_{EM} , T_{CM} and T_{EMRA} with the CCR7 and CD45RO markers. Amongst CD3⁺ T cells, effector memory T cells predominate in the human ocular surface. In earlier experiments, it has also been shown that CD8⁺ memory cells are the major populations in the conjunctival epithelial T cells among the healthy individuals²⁰. **Figure 3** shows that CD4⁺ and CD8⁺ effector memory T cells (T_{EM} and T_{EMRA}) are the major subset in the human ocular surface in 10 healthy controls studied. Each population mentioned in the figure are notified with their marker phenotype and names (Naïve, T_{CM} , T_{EMRA}). The numbers in red denote the percentage of the population. The data in this figure are represented as mean ± SEM

Impression cytology



Figure 1: Collection of IC Samples from the Ocular Surface by the Impression Cytology Device. Samples were collected from the temporal bulbar region as shown.



Figure 2: Dot Plot Graph using Flow Cytometry. Live 7-AAD⁻ cells were selected. The CD3⁺ cells were first gated, and then this population was gated into CD4⁺ and CD4⁻ subsets. The proportions of naïve (CCR7⁺CD45RO⁻), central memory (CCR7⁺CD45RO⁺) and effector memory (CCR7⁻CD45RO⁺; CCR7⁻CD45RO⁻) subsets was determined with the help of CCR7 and CD45RO markers. Please click here to view a larger version of this figure.



Figure 3: Immune Cell Sub-types in CD4⁺ and CD8⁺ T Cells of Healthy Human Ocular Surface. Distribution of conjunctival CD4⁺ and CD8⁺ naïve, central memory (T_{CM}), and effector memory (T_{EM} and T_{EMRA}) subsets in healthy human controls; each data point represents a separate individual mean ± SEM shown. The percentage of the population is marked as red below the name of each of the populations. The total percentage of the population is 98%, because CD4⁺ T_{EMRA} population was not included in the mentioned scatterplot (modified from Bose *et al.*²¹). Please click here to view a larger version of this figure.

Discussion

This is an easy, quick and less invasive technique that can be used in the out-patient clinics for relatively fast immune profiling in contrast to the conventional techniques like scraping, swabbing, pipetting or absorbent filter paper^{1,2}. A variant of this technique is already being used in research settings²². The future application of the proposed methodology is for patient stratification in clinical trials with ocular diseases, especially those requiring immunophenotyping.

A major challenge with this technique is the relatively few immune cells retrieved after impression collection and scraping. The total number of $CD3^+$ T cells recovered from four impressions per individual varied from ~ 500 - 1,000 cells. The ocular samples were washed before the flow cytometry analysis for a minimal number of times to avoid further loss of cells. Critical steps and challenges that remain within the protocol are efficient collection of ocular samples and proper scraping of the membrane to achieve higher cell numbers. Nevertheless, this limitation is unlikely to bias towards any specific immune phenotype. The troubleshooting performed here to maximize the yield of cells was to reduce the number of washing steps after and before the incubation of antibodies.

There are other limitations to using IC. In patients with severely keratinized or fibrosed ocular surface such as in Steven Johnson syndrome, the cellular yield may be even less than that in this study. The proportion of immune cells may change if the samples are stored instead of analyzed on the same day. It is difficult to predict if certain cell types are more resistant to storage than others. Previous studies have reported elevated levels of HLA-DR expression in the conjunctival epithelial cells²³, so it would be interesting to evaluate the correlation between HLA-DR and levels of specific immune cells. Immune cells may also be associated with the expression level of chemokines. These issues should be addressed in future studies.

Disclosures

The authors declare no competing financial interests.

Acknowledgements

The authors would like to thank Nandini Nallappan and Sharon Yeo for helping with the technical steps. The study was funded by a Start-Up Grant to KGC from Lee Kong Chian School of Medicine, Nanyang Technological University and by a Senior Clinician Scientist Award from the Singapore Ministry of Health's National Medical Research Council (NMRC) to LT (NMRC/ CSA/ 045/ 2012) and by a grant from the NMRC and administered by the National Health Innovation Centre to KGC and LT (NHIC-12D-1409007).

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