Stimulation of IL-2 Secretion in Human Lymphocytes
Using the Lionheart LX Automated Microscope to Image and Analyze Silver-stained ELISPOT Assays

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Abstract

Lymphocytes are used in several fields of research including autoimmune disorders, infectious diseases, vaccine development and hematological malignancies. The ELISPOT assay monitors ex-vivo cellular immune responses to antigenic stimuli. While the original ELISPOT assays used PVDF membrane plates, recently the assay format has developed to use a silver staining process that allows clear bottom plates to be utilized. Here we use the Lionheart™ LX Automated Microscope in conjunction with Gen5™ Microplate Reader and Imager Software to quantitate changes in IL-2 secretion in Jurkat cells using a silver-stain ELISPOT assay format.

Introduction

Cytokines are small molecular weight proteins or peptides secreted by many cell types (particularly immune system cells) that regulate the duration and intensity of the immune response. The cytokine interleukin 2 (IL-2) is a pleiotropic cellular regulatory molecule that is produced by lymphoid cells in response to several stimuli. It plays a role in preventing autoimmune diseases by promoting differentiation of immature T cells into regulatory T cells.[1] In addition, it causes the differentiation of T cells into effector T cells and memory T cells when the original T cell was stimulated by an antigen.[2]

T-cell activation is normally initiated by the interaction of a cell surface receptor to its specific ligand molecule along with a co-stimulatory molecule.[3] This binding event triggers the rapid hydrolysis of inositol phospholipids to diacylglycerol and inositol phosphates by phospholipase C (PLC). Diacylglycerol is an allosteric activator of protein kinase C (PKC) activation and inositol phosphates, which trigger Ca++ release and mobilization, resulting in a cascade of additional cellular responses mediating T-cell activation. One of these cellular responses is the production and secretion of IL-2. Triptolide is a diterpene triepoxide, that is a potent immunosuppressant and anti-inflammatory agent. Triptolide has been shown to inhibit the expression of IL-2 in activated T cells at the level of purine-box/nuclear factor and NF-κB mediated transcription activation. (Figure 1).[4]

While Jurkat cells are a leukemic T-cell line known to produce IL-2, under normal growth conditions little IL-2 is produced. Only after stimulation will substantial amounts of the cytokine be expressed.[5] Phytohemagglutinin (PHA) is a lectin that binds to the sugars on glycosylated surface proteins, including the T cell receptor (TCR), and nonspecifically binds them. The result is the low level stimulation of the signal cascade required for IL-2 secretion.[6] Likewise, Phorbol myristate acetate (PMA) is a small organic compound, which has a structure analogous to diacylglycerol, that diffuses through the cell membrane into the cytoplasm where it directly activates Protein Kinase C (PKC), Used in combination with Ionomycin, a calcium ionophore that triggers calcium release also results in producing a low level signal. However, when PMA and a co-stimulator, such as PHA, stimulate Jurkat cells IL-2 concurrently, production is strongly enhanced.[7]

The silver spot ELISPOT assay procedure is very similar to that of a conventional ELISPOT. The plates are first coated with the appropriate capture antibody. Cultured secreting cells

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Cytokine
Jurkat Cell
Brightfield
Silver Stain
IL-2
are added to the wells along with any experimental mitogen or antigen. Cells are maintained for a period of time after which they are removed. The analyte remains bound to the capture antibodies in close proximity to the location on the plate where the cell that secreted the analyte was situated. After removal of the cells and any unbound materials, a biotinylated detection antibody is added followed by an enzyme conjugate with incubation to allow binding and wash to remove unbound materials after each. Spots are formed with a substrate that deposits silver on the plate bottom at the locations of the original analyte capture. The resultant spots are then analyzed/ counted by image analysis (Figure 2).

Figure 2. ELISPOT silver stain Procedure.

The main difference between the ELISPOT assays used in this study is that the microplates use a transparent membrane, allowing inverted geometry microscopes to be used to quantify the silver spots. This allowed use of the Lionheart™ LX Automated Microscope.

Materials and Methods

Human IL-2 ELISPOT silver stain kit was obtained from U-CyTech biosciences (Utrecht, The Netherlands). Phorbol 12-myristate (PMA), cat # (P8139), and phytohemagglutinin-L (PHA), cat#11249738001 and Triptolide (cat # XXXX) were purchased from Millipore-Sigma. Ionomycin (cat# 407952) was from EMD-Millipore. Jurkat cells (human lymphoma) were from ATCC.

Cell Culture

Jurkat cells were grown in RPMI-1640 plus 10% FBS supplemented with 2 mM glutamine, penicillin and streptomycin. The suspension cells were split 1:5 with fresh media every 2-3 days. For experiments, unless otherwise indicated cells were plated at a density of $1 \times 10^5$/well in 96-well plates, previously coated with IL-2 antibody. Jurkat cells were stimulated to secrete IL-2 with a combination of PMA (50 ng/mL), Ionomycin (1 µg/mL) and PHA (1 µg/mL).

Triptolide Inhibition

Jurkat cells were plated at $1 \times 10^5$/well in 50 µL volume of complete RPMI media. After allowing cells to recover for 1 hour at 37 °C, in a humidified 5 % CO₂ environment, triptolide treatment was added in complete RPMI media at 4x of final concentration to each well in 50 µL. After treating with triptolide for 1 hour, IL-2 stimuli mixture (2x) was added in 100 µL for a final volume of 200 µL. After stimulating for 24 h, plates were assayed using an ELISPOT kit.

ELISPOT Assay

A Human IL-2 ELISPOT kit from U-CyTech Biosciences was used for these experiments. The assays were performed according to the kit instructions. Briefly, the plates are first coated with the appropriate concentration of capture antibody and allowed to absorb overnight at 4 °C. The unbound antibody is aspirated and the plate is manually washed 3x with PBS. The wells are then filled with a blocking solution (200 µL) and allowed to incubate for at least 1 hour at room temperature. Cultured Jurkat cells are added to the wells along with any experimental mitogen or inhibitor in a total volume of 200 µL. Cells are maintained for a period of 24 hours after which they are removed by washing 6x with 250 µL PBS-Tween 0.05% using a MultiFlo™ FX Multi-Mode Dispenser. A biotinylated detection antibody (100 µL) is added to the well and allowed to incubate for 60 minutes at 37 °C, after which unbound detection antibody is removed by washing. A streptavidin-GABA conjugate is then added (100 µL) and incubated at 37 °C for 60 minutes. Again, unbound conjugate is removed by washing and two-part substrate is then added that deposits silver grains onto the well bottom. Reactions are halted after 60 minutes at RT by washing with deionized water (250 µL) 3x and allowed to dry. Entire wells were then imaged.

Plate Washing

Plates were washed according to the assay kit instructions using a MultiFlo™ FX Washer Dispenser (BioTek Instruments). Wash buffer consisted of PBS (NaCl 137 mM, KCl 2.7 mM, Na₂HPO₄ 10 mM, KH₂PO₄ 7.4 mM) supplemented with 0.05% Tween 20. Unless specifically indicated, plates were washed six times with 250 µL buffer per well.
Plate Imaging

Cultures were imaged using a Lionheart™ LX Automated Microscope (BioTek Instruments, Winooski, VT) configured with enhanced brightfield. The imager uses a combination of LED light sources in conjunction with band pass filters and dichroic mirrors to provide appropriate wavelength light. A series of images were taken with a 4x objective to create a montage (4x5), that covered the entire well. Images were focused automatically using brightfield dual peak autofocus routine in Gen5™.

Analysis

Montage tiles (20 per well) were first stitched into a single image file prior to background subtraction. After background subtraction, objects were identified based on a threshold value and size criteria (Table 1).

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Table 1. Image processing and Image-analysis Parameters for ELISPOT Quantitation.

Results

Initial experiments demonstrate the specificity of the ELISPOT reaction. As seen in Figure 3, Jurkat cells that have been stimulated with a combination of PMA/Ionomycin and PHA exhibit numerous spots, while unstimulated cells produce few if any silver spots. Treatment alone without Jurkat cells also does not produce any spots.

![Figure 3. Specificity of IL-2 ELISPOT Reaction.](image)

The deposited silver from the conversion of the substrate by the conjugate enzyme results in very dark spots against a light background. It was noted that with wells with high numbers of spots that the intensity of the spot was diminished. The root cause of this is most likely depletion of the reagents within the well. In order to capture all of the spots under these circumstances a threshold value of 15000 was used for all experiments. The relationship to the threshold value and positive objects from several wells is depicted in Figure 4.

![Figure 4. Scatterplot of the Object Mean Intensity vs. Object number.](image)

Correct sizing of the identified objects is critical for accurate determinations. The intent of the ELISPOT assay is to identify and quantitate the number of cells responding to specific stimuli. The antibody-coated plate captures its specific target rather than the actual secretory cell. While most of the secreted analyte will be captured in the area immediately surrounding the position of the cell, some of the analyte will diffuse into the media and captured elsewhere. The high concentration of analyte near the cell will result in a spot as large or larger than the physical size of the cell, while dispersed analyte will result in
very small silver deposits. Because the density of the deposited silver grains is very high, it is critical that the selected spots are sized appropriately. Figure 5 demonstrates the number of dense silver grains 1 µm or greater present in a typical ELISPOT well. Only those spots exceeding 50 µm in size are designated as true spots.

Figure 5. Scatterplot of object size vs object number. All spots achieving a threshold of 15000 greater than 1 µm in size were plotted against their designation number. Spots designated as positive (size >50 µm) are indicated in RED.

The number of recorded spots produced from stimulated cells is proportional to the number of secreting cells. When a titration of Jurkat cells are exposed to a fixed concentration of stimulant the number of counted spots is proportional to the cell number. As demonstrated in Figure 6, increasing numbers of cells in a well results in an increase in the number of spots counted. Cell counts above 20,000 per well resulted in the spots coalescing together. Subsequent experiments used 10,000 cells per well.

Figure 6. Jurkat Cell Titration. Jurkat cells were seeded at various concentration into an ELISPOT plate and stimulated with 50 ng/mL PMA, 1 µg/mL Ionomycin, and 30 µg/mL PHA for 24 hours. The ELISPOT plate was then assayed for IL-2 secretion. Data points represent the mean of 8 determinations.

Secretion of IL-2 following stimulation of Jurkat cells is time dependent. Virtually no spots are observed in the first two hours after stimulation. Maximum secretion, as determined by spot number, was observed after approximately 8-10 hours in the presence of the stimulant. Further exposure did not result in an increase in the number of observed spots (Figure 7).

Figure 7. The Effect of Stimulation Exposure time on IL-2 secretion. Jurkat cells were exposed to 50 ng/mL PMA, 1 µg/mL Ionomycin and 30 µg/mL PHA for various periods of time before being assayed for IL-2 secretion using a silver staining ELISPOT assay. Data points represent the mean of 6 determinations.

Stimulation of IL-2 secretion by a mixture of PMA, Ionomycin and PHA is dose dependent. As observed in Figure 8, when a serial dilution of the compound mixture is used to stimulate IL-2 secretion, with increasing dilution (lower concentration) fewer spots are observed with the ELISPOT assay for IL-2.

Figure 8. Titration of Stimulate Mixture. Jurkat cells were stimulated with various dilutions of PMA/Ionomycin and PHA mixture for 24 hours in an ELISPOT plate coated with IL-2 antibody. Concentration is expressed as a percentage of the highest concentration (50 ng/mL PMA, 1 µg/mL Ionomycin, 30 µg/mL PHA). After stimulation IL-2 secretion was assayed and spots counted. Data points represent the mean of 6 determinations.
Pretreating Jurkat cells with Triptolide for 1 hour prior to stimulation reduces IL-2 secretion in a dose dependent manner. As demonstrated in Figure 9, increasing concentrations of triptolide result in fewer spots indicative of an IL-2 secreting cell. In these experiments, a stimulatory dose that was 80% of maximal was employed. The IC<sub>50</sub> under these conditions was determined to be 40 nM, which is similar to reports in the literature<sup>[8]</sup>.

**Discussion**

These data demonstrate the utility of the Lionheart™ LX Automated Microscope in conjunction with Gen5™ Microplate Reader and Imager Software to image and analyze silver stained ELISPOT assay plates. The combination of a PHA/Ionomycin and PHA have been shown to markedly stimulate IL-2 secretion in cultured Jurkat cells. Without stimulation, IL-2 is virtually absent. The stimulation of IL-2 secretion is both dose and time dependent. The ability of Triptolide, a known transcription inhibitor, to prevent IL-2 secretion suggests that new protein synthesis is required after stimulation.

ELISPOT is a sensitive assay to monitor the ex vivo cellular immune response at the single cell level by detecting secreted proteins released by cells. This technique has been derived from the sandwich enzyme-linked immunosorbent assay (ELISA) to accommodate the use of whole cells to identify the frequency of the secreting cells. As such, there are a number of critical parameters that need to be optimized in order for experiments to be successful. Depending on the degree of cellular secretion, developed spots can be quite large. The expected number of positive cells is of greater importance than the total number of cells used initially. The presence of too many secreting cells results in the individual spots coalescing making a numerical determination difficult. For example, an investigation of a relatively rare secreting event would require a greater number of cells to be seeded as compared to a more common event. Timing of the response relative to the stimulation and/or the inhibition is important. Receptor mediated events often will take longer to elicit a response than a stimulatory molecule that can interact within the cell directly. It is important that an appropriate interval between stimulation and measurement be utilized. The testing of inhibitors still requires a stimulating agent to be present. In these experiments, it is important that a less than maximal concentration of the stimulatory agent be used, lest it mask any inhibitory affects.

The Lionheart LX is an ideal platform to interpret silver stained ELISPOT assays. The imager supports microscopic imaging with objectives from 1.25x to 100x that can be installed in a 6-position objective turret. While this research only used brightfield illumination, the imager supports fluorescence based microscopy with LED and filter cubes. Gen5 Microplate Reader and Imager Software, besides controlling reader function, can be used to automatically perform stitching of separate montage image tiles, perform background subtraction and mask off regions outside the well prior to analysis.

**References**


