NCL Method ITA-33

Detection of Changes in Mitochondrial Membrane Potential in T-cells Using JC-1 Dye

This protocol assumes an intermediate level of scientific competency with regard to techniques, instrumentation, and safety procedures. Rudimentary assay details have been omitted for the sake of brevity.
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1. Introduction

The purpose of this protocol is to understand the propensity of nanomaterials to induce mitochondrial damage as assessed by change in mitochondrial membrane potential (ΔΦm). Alteration in mitochondrial physiology is commonly linked with apoptotic cell death. Under stressed conditions, the mitochondrial permeability transition pores open, leading to a decrease in mitochondrial membrane potential [1]. The changes in ΔΦm further activates various signaling cascades leading to cytochrome C release, ATP depletion and accumulation of reactive oxygen species (ROS), leading to cell death via either necrosis or apoptosis [1, 2]. The combination of mitochondrial membrane potential loss and mitochondrial ROS lead to organelle specific toxicity that ultimately cause cell death.

2. Principles

T-lymphocytes are isolated from human blood anti-coagulated with Li-heparin using RosetteSep™ reagent from STEMCELL Technologies, followed by gradient separation using Ficoll-Paque Plus solution. The blood is obtained from three healthy donor volunteers. Specimens from each donor are processed separately. Isolated T-cells are incubated with test-nanoparticles, and the changes in mitochondrial membrane potential (ΔΦm) are detected using JC-1 reagent. JC-1 is a cell-permeable fluorogenic probe which fluoresces in Red (FL-2) when present in aggregates, but fluoresces in green (FL-1) when present in monomer (Figure 1).

Note: Cell lines/PBMC can be used instead of T-cells.

![Figure 1. Principle of JC-1 Reagent.](image)

(A) Healthy mitochondria accumulates JC-1 inside mitochondrial cristae. (B) Damaged mitochondria accumulates JC-1 monomers outside mitochondria. (C) The transition of JC-1 dye form aggregates to monomer shows different fluorescence.
3. Reagents, Materials, and Equipment

Note: The NCL does not endorse any of suppliers listed below; their inclusion is for informational purpose only. Equivalent supplies from alternative vendors can be substituted.

3.1 Reagents

1. Phosphate buffer saline (PBS) (HyClone, AQB 22934)
2. Ficoll-Paque Premium (GE HealthCare, 17-5442-03)
3. Fetal bovine serum (HyClone, SH30070.03)
4. RPMI1640 (Invitrogen, 11875-119)
5. Pen/Strep solution (Invitrogen, 15140-148)
6. L-glutamine (HyClone, SH30034.01)
7. Cellometer ViaStain AOPI Staining Solution in PBS (Nexcelom Biosciences, CS2-0106; or other reagent for distinguishing between live and dead cells)
8. Albumin, from bovine serum (BSA) (Sigma, A4503)
9. MitoProbe™ JC-1 Assay Kit (Molecular Probes, M34152)
10. Hank’s balanced salt solution (HBSS) (Invitrogen, 24020-117)
11. Dimethyl sulfoxide (DMSO) (Sigma, D2650)
12. RosetteSep™ (STEMCELL technologies, 15061)
13. tert-butyl hydroperoxide (TBHP) (Acros organics, AC180340050)

3.2 Materials

1. Pipettes, 0.05 to 10 mL
2. Polypropylene tubes, 50 and 15mL
3. 24/48 well cell culture plates (Corning, 3524, 3548)
4. Polystyrene round bottom 12 x 75 mm² tubes (Falcon tubes, 352058)

3.3 Equipment

1. Refrigerator, 2-8°C
2. Freezer, -20°C
3. Cell culture incubator with 5% CO2 and 95% humidity
4. Biohazard safety cabinet approved for level II handling of biological material
5. Vortex
6. Cellometer, Source or Hemacytometer to perform cell count
7. Flow Cytometer (e.g., FACSCalibur)
8. Centrifuge, 1200xg

4. **Reagent and Control Preparation**

4.1 Complete RPMI

The complete RPMI medium should contain 10% FBS (heat-inactivated), 4 mM L-glutamine, 100 U/mL penicillin, and 100 µg/mL streptomycin sulphate. Store at 2-8°C protected from light for no longer than 1 month. Before use, warm the medium in a 37°C water bath.

4.2 Heat-inactivated fetal bovine serum

Thaw a bottle of FBS at room temperature or overnight at 2-8°C and allow to equilibrate to room temperature. Incubate 30 minutes in a 56°C water bath, mixing every 5 minutes to heat-inactivate it. Single-use aliquots may be stored at 2-8°C for up to one month or at a nominal temperature of -20°C indefinitely.

4.3 Positive Control (CCCP)

The kit provides 125 µL of 50 mM CCCP dissolved in DMSO. Add 1 µL of CCCP to appropriate wells during staining protocol and incubate the sample at 37ºC for 5 min.

4.4 Negative Control

Use PBS as a negative control.

4.5 JC-1 Reagent

Briefly, centrifuge the JC-1 vial before opening. The kit provides 30 µg powdered dye. Prepare 200 µM intermediate dilution by adding 230 µL of DMSO to the vial. Add 10 µL of the intermediate JC-1 dilution to 1 mL of cell suspension. Alternatively, use 300 µL of medium with 2 µM JC-1 per sample and incubate for 15-30 min at 37°C.

4.6 Vehicle Control

Vehicle control is the buffer or media used to formulate test nanomaterials. Common excipients used are trehalose, sucrose, and albumin. However, other reagents and materials are also used alone or in combination. Vehicle control should match formulation buffer of the test-nanomaterial by both composition and concentration. This control can be skipped if nanoparticles are stored in PBS.
5. Preparation of Nanoparticles

When the experiment is conducted in 24-well plates, the assay requires 1.5 mL of nanoparticles dissolved/re-suspended in complete culture medium at a concentration 5X higher than the highest final test concentration. Test concentrations are based on the calculated plasma concentration of the nanoparticle at the intended therapeutic dose. For the purpose of this protocol this concentration is called “theoretical plasma concentration”. Considerations for estimating theoretical plasma concentration have been reviewed elsewhere [3] and are summarised in Box 1 below.

This assay will evaluate four concentrations: 10X (5X if 10X cannot be achieved, or 100X or 30X when feasible) of the theoretical plasma concentration, the theoretical plasma concentration, and two 1:5 serial dilutions of the theoretical plasma concentration. When the intended therapeutic concentration is unknown, 1 mg/mL is used as the highest concentration. Alternatively, the highest reasonably achieved concentration can be used if 1 mg/mL is unattainable.

For example, if the theoretical plasma concentration to be tested is 0.2 mg/mL, a stock of 10 mg/mL is prepared. This sample is then diluted 10-fold (1.0 mg/mL), followed by two 1:5 serial dilutions (0.2 and 0.04 mg/mL). When 0.2 mL of each of these sample dilutions are combined in a culture plate with 800 µL cells, the final concentrations of nanoparticles are 0.008, 0.04, 0.2, and 2 mg/mL. Each nanoparticle concentration is plated two times.

Box 1. Example Calculation to Determine Nanoparticle Concentration for In Vitro Tests

In this example, we assume a mouse dose of 123 mg/kg. Therefore, the scaled equivalent human dose would be:

\[
\text{Human dose} = \frac{\text{mouse dose}}{12.3} = \frac{123 \text{ mg/kg}}{12.3} = 10 \text{ mg/kg}
\]

Blood volume constitutes approximately 8% of the body weight. Therefore, an average human of 70 kg body weight has approximately 5.6 L of blood. Assuming all the nanoparticle injected goes into the systemic circulation, this provides a rough approximation of the potential maximum nanoparticle concentration in blood, which is used as the in vitro test concentration.

\[
\text{in vitro concentration}_{\text{human matrix}} = \frac{\text{human dose}}{\text{human blood volume}} = \frac{70 \text{ kg} \cdot 10 \text{ mg/kg}}{5.6 \text{ L}} = 0.125 \text{ mg/mL}
\]
6. Collection and Handling of Whole Blood

Collect whole blood from three healthy donor volunteers who have been not on medication and who have been clear from infection for at least 2-weeks before blood donation. Use Li-heparin tubes and discard the first 10 cc. For best results, whole blood should be used within 1 hour after collection. Prolonged storage (> 2 hr) of whole blood will lead to a decrease in cell function.

7. Preparation of T-Cells

7.1. Place freshly drawn blood into 15 or 50 mL conical centrifuge tubes. Add 50μL RosetteSep per mL of blood, mix well, and incubate the tubes at room-temperature for 20 minutes.

7.2. Add equal amount of room-temperature PBS with 2% FBS to blood and mix well.

7.3. Slowly layer the Ficoll-Paque solution underneath the blood/PBS mixture by placing the tip of the pipet containing Ficoll-Paque at the bottom of the blood sample tube. Alternatively, the blood/PBS mixture may be slowly layered over the Ficoll-Paque solution. Use 1 mL of Ficoll-Paque solution per 2 mL of blood/PBS mixture. For example, 15 mL Ficoll-Paque is required for 15 mL blood diluted with 15 mL PBS with 2% FBS in a 50 mL tube.

Note: To maintain Ficoll-blood interface it is helpful to hold the tube at a 45º angle.

7.4. Centrifuge 20 min at 1200xg, 18-20ºC, without brake.

Note: Depending on the type of centrifuge, one also may need to set acceleration speed to minimum.

7.5. Using a sterile pipet, remove the upper layer containing plasma and platelets and discard it.

7.6. Using a fresh sterile pipet, transfer the T-lymphocyte cell layer into another centrifuge tube.

7.7. Wash cells by adding an excess of HBSS and centrifuging for 10 min at 400xg, 18-20ºC. The HBSS volume should be ~3 times the volume of the mononuclear layer.

Note: Usually 20 mL of blood/PBS mixture results in ~ 1 mL of the T-cell layer and requires at least 3 mL of HBSS for the wash step. We use 5 mL of HBSS per each 1 mL of cells.
7.8. Discard supernatant and repeat wash step once more.

7.9. Re-suspend cells in complete RPMI-1640 medium. Dilute cells 1:2 with AOPI dye and load 20 µL in Cellometer slide. Report cell count and viability displayed by Cellometer. If viability is at least 90%, proceed to step 8.

*Note: Similar assay can be applied for the PBMC isolated according to ITA-10.*

8. **Treatment with Nanoparticles and Controls**

8.1. Adjust cell count to 1x10⁶ cells/mL.

8.2. Aliquot 400 µL of cell suspension prepared in step 8.1 per well of 48-well plate.

8.3. Add 100 µL of test-nanoparticles and control to appropriate wells and incubate for appropriate length of the assay at 37°C/5% CO₂. (Add 100 µL of complete media to “unlabeled cells” wells).

*Note: For the positive control inhibition, it is essential to perform PC treatment immediately after NC treatment for an additional 30 minutes. Longer incubation after NC treatment is ineffective when tested for inhibition on PC.*

8.4. Wash cells once with 3 mL of PBS.

8.5. Add the JC-1 reagent to all samples except “unlabeled cells” and incubate for 30 minutes at 37°C/5% CO₂.

8.6. Wash cells once with 3 mL of PBS.

8.7. Reconstitute cells in 300 µL PBS and analyze samples by flow cytometry.

*Note: It is critical to run samples on flow cytometer within 15 min after staining, as longer incubation may result in release of JC-1 aggregates from healthy mitochondria and may fail acceptance criteria for NC staining.*

9. **Flow Cytometry**

*At the NCL, we use FACSCalibur. The procedure below is based on our experience with this instrument. If you are using another cytometer, please follow the procedure specific to that instrument.*

9.1. Switch on the instrument. Make sure the sheath tank is full and the waste tank is empty.

9.2. To get consistent results, run BD FACSComp software using BD CaliBRITE beads.
9.3. Adjust FSC vs. SSC dot plot using unlabeled cells to get the population of cells in the plot. See Figure 2 below for example.

9.4. Set the threshold to remove most of the debris.

9.5. Gate lymphocyte population. See Figure 2 below for example.

   Note: The purified T cells do not need to be gated unless the cells show multiple dense clusters on the FSC vs. SSC plots.

9.6. Create FL2 vs. FL1 histogram for each analyzed cell population. See Figure 3 for example.

9.7. Create Gate 2 for FL2 positive cells using CTRL sample.

9.8. Create Gate 3 for FL1 positive cells using CCCP sample

9.9. Analyze shift of % population in Gate 3 from Gate 2.

9.10. Acquire and save the data.

9.11. Follow the instrument closing procedure.

**Figure 2.** Set up gates for lymphocytes: gate 1.

**Figure 3.** Gating of lymphocytes containing JC-1 aggregates (gate 2) and JC-1 monomer (gate 3) using NC and PC samples, respectively, in FL2 vs. FL1 dot plot. (Left) untreated control sample stained with JC-1 reagent. (Right) CCCP treated sample stained with JC-1 reagent. (blue–NC, purple–PC).
10. Data Analysis and Report

Use appropriate software to analyze the data acquired in step 9.10. CellQuest or other flow cytometry software can be used. Here, FCSEExpress from De Novo Software Solutions, Inc. was used. Report the data as % positive cells in FL-1 indicating cells with damaged mitochondria. Alternatively, ratio of MFI in FL2/FL1 can be used as an assessment of healthy/damaged mitochondria in each treatment group.

11. Acceptance criteria

11.1 The run is acceptable if PC shows more than 80% cells in FL1, and NC shows more than 70% in FL2.

12. References


13. Abbreviations

DMSO – dimethyl sulfoxide
FL - fluorescence
HBSS – Hank’s balanced salt solution
NAC – N-acetyl-L-cysteine
NC – negative control
PBS – phosphate buffered saline
PC – positive control
RPMI – Roswell Park Memorial Institute
TBHP – tert-butyl hydroperoxide
**TS** – test samples

**VC** – vehicle control

### 14. Appendix

**Example Plate Map**

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<tr>
<td>A</td>
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<td>VC</td>
<td>PC</td>
<td>PC + NAC</td>
<td>TS (0.008 mg/mL)</td>
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<td>B</td>
<td>TS (0.04 mg/mL)</td>
<td>TS + NAC (0.04 mg/mL)</td>
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**NC** – negative control, **VC** – vehicle control, **PC** – positive control, **NAC** – N-acetyl-L-cysteine, **TS** – test sample; shaded cells – inhibition control