Analysis of Apoptosis and Necroptosis by Fluorescence-Activated Cell Sorting

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Fluorescence-activated cell sorting (FACS) is a laser-based, biophysical technology that allows simultaneous multiparametric analysis. For the analysis of dying cells, fluorescently labeled Annexin V (Annexin V<sub>FITC</sub>) and propidium iodide (PI) are the most commonly used reagents. Instead of PI, 4',6-diamidino-2-phenylindole (DAPI) can also be used. DAPI is a fluorescent stain that binds strongly to A-T-rich regions in DNA. DAPI and PI only inefficiently pass through an intact cell membrane and, therefore, preferentially stain dead cells. DAPI can be combined with Annexin V<sub>FITC</sub> and the potentiometric fluorescent dye, tetramethylrhodamine methyl ester (TMRM), which measures mitochondrial permeability transition and mitochondrial membrane depolarization. TMRM is a cell-permeable fluorescent dye that is sequestered to active mitochondria, and hence labels live cells. On apoptosis or necroptosis the TMRM signal is lost. The advantage of using Annexin V<sub>FITC</sub>/DAPI/TMRM is that the entire cell population is labeled, and it is easy to distinguish living (TMRM+/Annexin V<sub>FITC</sub>+/DAPI-), from dying or dead cells (apoptosis: TMRM-/Annexin V<sub>FITC</sub>+/DAPI-; necrosis: TMRM-/Annexin V<sub>FITC</sub>+/DAPI+). This is important because cell debris (fluorescent negative particles) must be avoided to establish the correct parameters for the FACS analysis, otherwise incorrect statistical values will be obtained. To obtain information on the cell concentration or absolute cell counts in a sample, it is recommended to add an internal microsphere counting standard to the flow cytometric sample. This protocol describes the FACS analysis of cell death in HT1080 and L929 cells, but it can be readily adapted to other cell types of interest.

MATERIALS

It is essential that you consult the appropriate Material Safety Data Sheets and your institution’s Environmental Health and Safety Office for proper handling of equipment and hazardous materials used in this protocol.

RECIPIES: Please see the end of this protocol for recipes indicated by <R>. Additional recipes can be found online at http://cshprotocols.cshlp.org/site/recipes.

Reagents

4',6-Diamidino-2-phenylindole (DAPI)
Annexin V<sub>FITC</sub>
CaCl<sub>2</sub> (1 M)
CountBright Absolute Counting Beads (Life Technologies)
Doxycycline

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Dulbecco’s modified Eagle’s medium (DMEM) with phenol red (Life Technologies, 41966-029)
Fetal bovine serum (FBS)
HT1080indRIPK3 cells
  Parental HT1080 cells (ATCC) were transduced with lentiviral particles carrying RIPK3 cDNA expressed under
  the control of a doxycycline regulatable promoter. For RIPK3 induction, cells should be cultured in media
  containing doxycycline (DOX) (100 ng/mL) for at least 1 wk prior to the experiment.
IAP-antagonist (SMAC mimetic, SM; 1 mM in DMSO)
L929 cells
Necrostatin-1 (Sigma-Aldrich; 40 mM in DMSO)
Recombinant human TNF (Alexis; 100 µg/mL in DMEM)
Tetramethylrhodamine methyl ester (TMRM) (Life technologies)
Trypsin
zVAD-fmk (20 mM in DMSO)

**Equipment**

Cell culture plates (six-well)
BD LSR II FACS system (BD Bioscience) or equivalent
  The system needs to be equipped with three lasers: 404, 488, and 561 nm. FITC is excited by the 488 nm laser
  and emission light is measured with filters 505 LP 525/50 BP. DAPI is excited by the 404 nm laser and emission
  light is collected with filters 450/50 BP (Kapuscinski 1995). TMRM is excited by a 561 nm laser and emission light
  is collected with filters 595/40 BP. The CountBright absolute counting beads can be excited by any laser and
  detected by any filter.
Incubator (37°C, 10% CO2, humidified)
Polystyrene round-bottom FACS tubes (5 mL)
Tissue culture equipment, including a laminar flow hood

**METHOD**

Perform Steps 1–3 under a laminar flow hood using aseptic techniques.

1. Seed 3 × 10^5 cells/well (HT1080indRIPK3 or L929) in 2 mL of DMEM supplemented with 10% FBS
   into six-well plates. Culture cells at 37°C under a 10% CO2 atmosphere.
2. Once the cells have reached 50%–60% confluency, treat them as follows. Ensure sufficient wells
   are left untreated for use as controls, as described in Step 6.
   i. Treat wells of HT1080indRIPK3 cells for 12 h with the following combinations of reagents. Use three wells per condition.
      • 20 µM zVAD-fmk
      • 10 ng/mL TNF and 500 nM SM
      • 10 ng/mL TNF and 500 nM SM and 20 µM zVAD-fmk
      • 10 ng/mL TNF and 500 nM SM and 20 µM zVAD-fmk and 40 µM necrostatin-1
   ii. Treat wells of L929 cells for 2 h with the following combinations of reagents. Use three wells per condition.
      • 10 ng/mL TNF + 500 nM SM + 20 µM zVAD-fmk
      • 10 ng/mL TNF + 500 nM SM + 20 µM zVAD-fmk + 40 µM necrostatin-1

   Cells can be supplemented with other drugs or treatment combinations appropriate for the experimental
   goals. For treatments up to 12 h there is no need to change the medium.
3. Aspirate the supernatants into prelabeled FACS tubes.
   This allows the collection of detached dead cells.
4. Trypsinize cells and carefully resuspend them with their own medium (collected in Step 3). Return each cell suspension to its respective FACS tube.

5. Add 2.5 mM CaCl$_2$, 5 µL/mL Annexin V$^{FITC}$, 1.43 µM DAPI, 20 nM TMRM, and CountBright beads (25000/mL), and proceed to Step 7.

   CountBright absolute counting beads are a calibrated suspension of microspheres that are brightly fluorescent across a wide range of excitation and emission wavelengths, and contain a known concentration of microspheres. For absolute counts, a specific volume of microsphere suspension is added to a specific volume of sample, so that the ratio of sample volume to microsphere volume is known. The volume of sample analyzed can be calculated from the number of microspheres counted, and can be used to correct for cell loss during analysis and to determine cell concentration.

6. Prepare controls as follows.
   i. Leave one well of HT1080$^{indRIPK3}$ cells and one well of L929 cells without any dye.
   ii. Treat HT1080$^{indRIPK3}$ and L929 cells with each of the following single dyes. Use one well of cells per dye.
      - 5 µL/mL Annexin V$^{FITC}$
      - 1.43 µM DAPI
      - 20 nM TMRM
      - CountBright beads (25,000/mL)

7. Incubate the cells from Steps 5 and 6 for at least 20 min at 37°C in the incubator to allow the dyes to adhere. 
   
   After this, cells can be kept on ice.

8. Analyze samples using an LSRII FACS system, or equivalent.
   i. Analyze the unstained control cells first and adjust the voltage and threshold.
   ii. Analyze the single-stained controls, and set up a compensation matrix.
   iii. Use the nonstained and single-stained control samples for gating. 

      The exact parameters and conditions for sorting will vary among FACS systems and facilities and should be discussed with experts in your facility.

9. Use three technical replicates per experimental condition and three biologically independent experiments for statistical analysis using Student’s $t$-test.

   For sample results, see Figure 1.

DISCUSSION

FACS provides a fast, objective, and quantitative method of recording of the number of dying cells in a population and is, therefore, routinely used to study cell death (Christensen et al. 2013). Annexin V is used as a probe to detect cells in which phosphatidylserine (PS) is exposed at the outer leaflet of the plasma membrane (Tait et al. 1989; Andree et al. 1990; Vermes et al. 1995). PS externalization occurs in early apoptotic cells, whereas living cells remain Annexin V negative. Although Annexin V can be used to detect apoptosis, it should be noted that necroptotic cells also become Annexin V-positive as Annexin V can bind to internal PS following cell rupture. However, when combined with PI the double-labeling procedure allows a further distinction of necrotic (Annexin V+/PI+) from early apoptotic (Annexin V+/PI$^-$) cells (Vermes et al. 1995). Although cells that are Annexin V+/PI$^-$ can be considered to die by apoptosis at early time points, it should be noted that at late time points apoptotic cells can become Annexin V+/PI+. Therefore, it is important to combine this assay with a time-course study, and/or the use of specific inhibitors. Although the FACS method is a powerful approach to study cell death, it is not without its caveats. A particular problem is the fact that dying cells can disintegrate and the resulting cell debris cannot be captured by FACS analysis. To circumvent
**FIGURE 1.** (See following page for legend.)

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FIGURE 1. FACS analysis of cells undergoing necroptosis. (A, B) HT1080ΔRIPK3 cells were grown in the absence or presence of 100 ng/mL Dox to induce RIPK3 expression. Cells were seeded into six-well plates and cultured for 24 h before being treated with 20 µM zVAD-fmk, 10 ng/mL TNF + 500 nM SM, or 10 ng/mL TNF + 500 nM SM + 20 µM zVAD-fmk. Prior to flow cytometry, cells were trypsinized and incubated with 5 µL/mL Annexin V-FITC, 1.43 µM DAPI, 20 nM TMRM and CountBright™ absolute counting beads (25,000/mL). (B) Graphic representation of cell death from (A) expressed as % of Annexin V-FITC and DAPI-positive (dead) cells, and corrected for the loss of cells using the microsphere counting beads. (C, D) L929 cells were seeded into 6-well plates for 24 h before being treated for 2 h with control, 10 ng/mL TNF + 500 nM SM + 20 µM zVAD-fmk, or 10 ng/mL TNF + 500 nM SM + 20 µM zVAD-fmk + 40 µM necrostatin-1. Prior to flow cytometry, cells were trypsinized and incubated with 5 µL/mL Annexin V-FITC, 1.43 µM DAPI, 20 nM TMRM and beads (25,000/mL). (D) Graphic representation of cell death expressed as a % of Annexin V-FITC and DAPI-positive (dead) cells, and corrected for the loss of cells using the microsphere counting beads.
this problem, CountBright absolute counting beads can be used to determine how many cells were lost due to cell disintegration. When using microspheres, it is important to take into consideration that the untreated controls will continue to proliferate, which might affect the cell count. Hence, the respective treatment times should not be too long. Despite its advantages and ease of use, FACS is best complemented with time-lapse video microscopy as the combination of both methodologies more accurately captures the modality of cell death. This is especially true for some cell lines that stain poorly for Annexin V when undergoing bona fide apoptosis (Fadeel et al. 1999; Lee et al. 2013). For example, PS exposure reportedly does not occur in autophagy-deficient cells succumbing to apoptosis (Qu et al. 2007). Nevertheless, such cells clearly lose cell mass through membrane blebbing and nuclear shrinkage, and at later stages they undergo secondary necrosis and become PI positive.

RELATED TECHNIQUES

In addition to the method described here, there are many other FACS-based assays that can be used, such as evaluation of the percentage of sub-G0 cells after permeabilization followed by PI or 7-aminoactinomycin D (7-AAD) staining (Tenev et al. 2001; Zembruski et al. 2012).

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REFERENCES


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