IncuCyte® Cell Proliferation Assay

For label-free proliferation measurements of adherent or non-adherent cell lines

This protocol provides an overview of the IncuCyte® Cell Proliferation Assay methodology. It is compatible with the IncuCyte® live-cell analysis system using your choice of cells and treatments. The highly flexible assay format can be combined with our range of IncuCyte® cell health and viability reagents for multiplexed measurements of cytotoxicity and apoptosis alongside proliferation in the same well.

Required materials

- Flat bottom tissue culture plate (e.g., Corning 3595)
- Poly-L-ornithine (Sigma P4957) – optional, for non-adherent cells
- Fibronectin (Sigma A7906) – optional, for non-adherent cells

General Guidelines

- Following cell seeding, place plates at ambient temperature (15 minutes for adherent cell lines and 45 minutes for non-adherent cell lines) to ensure homogenous cell settling.

- Remove bubbles from all wells by gently squeezing a wash bottle containing 70-100% ethanol with the inner straw removed, to blow vapor over the surface of each well.

- After placing the plate in the IncuCyte live-cell analysis system, allow the plate to warm to 37 °C for 30 minutes prior to scanning.

Quick Guide

1. COAT WELLS
   - Coat wells of plate (50 μL/well) with appropriate matrix. Optional for adherent cell lines.

2. PLATE CELLS
   - Seed cells (100 μL/well, 1,000–10,000 for adherent and 5,000–50,000 for non-adherent) into a 96-well plate.

3. ADD TREATMENTS
   - Add desired treatments (100 μL/well, 1x for adherent cell lines, 2x for non-adherent cell lines).
Adherent Cell Line Protocol

Day 0

1 Coat wells (optional)
   1.1 Depending on cell line used, coat a 96-well flat bottom plate with relevant coating matrix according to manufacturer’s recommendation.
   1.2 Prior to cell seeding, prepare cell treatments at 2x final assay concentration in enough cell culture medium to achieve a volume of 100 μL per well.

Plate Cells

2.1 Seed your choice of cells (100 μL per well) at an appropriate density into a 96-well plate, such that by day 1 the cell confluence is approximately 10 - 20%. The seeding density will need to be optimized for the cell line used; however, we have found that 1,000 to 2,500 cells per well (10,000 to 25,000 cells/mL seeding stock) are reasonable starting points.

Day 1

Add Treatments

1.1 Depending on cell line used, coat a 96-well flat bottom plate with relevant coating matrix according to manufacturer’s recommendation.
1.2 Prior to cell seeding, prepare cell treatments at 2x final assay concentration in enough cell culture medium to achieve a volume of 100 μL per well.

Plate Cells

2.1 Seed your choice of cells (100 μL per well) at an appropriate density into a 96-well plate, such that by day 1 the cell confluence is approximately 10 - 20%. The seeding density will need to be optimized for the cell line used; however, we have found that 1,000 to 2,500 cells per well (10,000 to 25,000 cells/mL seeding stock) are reasonable starting points.

Non-Adherent Cell Line Protocol

For label-free proliferation measurements of adherent or non-adherent cell lines

Day 1

Seed cells and add prepared treatments

1.1 Coat a 96-well flat bottom plate with relevant coating matrix. We recommend coating with 50 μL of either 0.01% poly-L-ornithine solution or 5 μg/mL fibronectin diluted in 0.1% BSA. Coat plates for 1 hour at ambient temperature, remove solution from wells, then allow plates to dry for 30-60 minutes prior to cell addition.
1.2 Prior to cell seeding, prepare cell treatments at 2x final assay concentration in enough cell culture medium to achieve a volume of 100 μL per well.

Plate Cells

2.1 Seed your choice of cells (100 μL per well) at an appropriate density into a 96-well plate. The seeding density will need to be optimized for the cell line used; however, we have found that 5,000 to 50,000 cells per well (50,000 – 500,000 cells/mL seeding stock) are reasonable starting points.

NOTE: If studying immune cell clustering and proliferation, prepare cell activation treatments at 5x final concentration, and immediately add 50 μL per well containing cells. It is advised that some control wells containing only vehicle are included in the plate.

Add Treatments

3.1 Immediately after cell seeding, add treatments and controls to appropriate wells of the 96-well plate containing cells. Triturate wells to appropriately mix the treatment to ensure cell exposure at 1x.
3.2 Place the cell plate into the IncuCyte live-cell analysis system and allow the plate to warm to 37°C for 30 minutes prior to scanning.
   a. Objective: 4x, 10x or 20x
   b. Channel selection: Phase Contrast (+ “Green” or “Red” if fluorescent label or cell health reagents are used)
   c. Scan type: Standard
   d. Scan interval: Typically, every 1 to 2 hours, until your experiment is complete.
Related Products and Applications

A comprehensive range of fluorescent nuclear labeling and cell health reagents are available for use with the IncuCyte Live-Cell Analysis System to enable multiplexed measurements of cytotoxicity and proliferation alongside apoptosis.

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<tr>
<th>Product</th>
<th>Cat No.</th>
<th>Amount</th>
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<td>IncuCyte® NucLight Red BacMam 3.0 Reagent for nuclear labeling</td>
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<td>IncuCyte® Cytotox Red Reagent for counting dead cells</td>
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<td>IncuCyte® Annexin V Red Reagent for apoptosis</td>
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<td>IncuCyte® Annexin V Green Reagent for apoptosis</td>
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A complete suite of cell health applications is available to fit your experimental needs. Find more information at essenbioscience.com

For additional product or technical information, please e-mail us at AskAScientist@essenbio.com
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