Product Manual

Radius™ 96-Well Cell Migration Assay

Catalog Number

CBA-126	96 assays
CBA-126-5	5 x 96 assays

FOR RESEARCH USE ONLY Not for use in diagnostic procedures



Introduction

Cell migration is a highly integrated, multistep process that orchestrates embryonic morphogenesis, tissue repair and regeneration. It plays a pivotal role in the disease progression of cancer, mental retardation, atherosclerosis, and arthritis. The initial response of a cell to a migration-promoting agent is to polarize and extend protrusions in the direction of the attractant; these protrusions can consist of large, broad lamellipodia or spike-like filopodia. In either case, these protrusions are driven by actin polymerization and can be stabilized by extracellular matrix (ECM) adhesion or cell-cell interactions (via transmembrane receptors).

The RadiusTM Cell Migration Assay Kit utilizes a proprietary 96-well plate to monitor the migratory properties of cells. Each plate well contains a 0.68 mm non-toxic, biocompatible hydrogel spot (RadiusTM Gel) where cells **cannot** attach. When adherent cells are seeded in the RadiusTM Cell Migration well, they attach outside of the RadiusTM Gel coated area. Once firm cell attachment is achieved, the hydrogel is quickly removed to expose a cell-free region to study cell migration/closure. This format provides a robust *in vitro* system to measure 2-D cell migration, screen potential inhibitors and study cytoskeleton reorganization events.

Additional features of the Radius[™] Cell Migration Assay:

- Exclusive coating method which produces consistent gel spot size 0.68 mm diameter ± 0.014 mm (2%)
- Qualitative, quantitative, real-time or endpoint analysis
- Radius[™] Cell Migration Plate does <u>not</u> need to be used all at once unused wells can be used in future experiments, up to a total of 3 migration experiment cycles
- Compatible with all cell stains, dyes, and labels
- Complete migration zone closure achievable in 15-30 hours (actual time is cell line dependent)
- Analyze by phase contrast or fluorescence microscopy
- Compatible with HCS/HCI instrumentation
- Adaptable to liquid handling equipment
- Optimized for 10X magnification (entire Radius[™] Gel migration area is viewed in a 10X magnification field)
- Radius[™] Gel removal is controlled and extremely fast (migration starts almost simultaneously between wells)

The RadiusTM Cell Migration Assay Kit is designed for High Content Analysis applications or imaging software and is adaptable to liquid handling equipment. Each kit provides sufficient quantities to perform 96 migration tests.



Assay Principle





Related Products

- 1. CBA-100: CytoSelectTM 24-Well Cell Migration Assay (Boyden Chamber Assay, Colorimetric)
- 2. CBA-101: CytoSelect[™] 24-Well Cell Migration Assay (Boyden Chamber Assay, Fluorometric)
- 3. CBA-110: CytoSelect[™] 24-Well Cell Invasion Assay (Boyden Chamber Assay, Colorimetric)
- 4. CBA-120: CytoSelect[™] 24-Well Wound Healing Assay
- 5. CBA-130: CytoSelect[™] 96-Well Cell Transformation Assay (Soft Agar Colony Formation)

RadiusTM 96-well Cell Migration Plate Dimensions

Well Volume	360 µL
Well Depth	10.67 mm
Well Diameter - Top	6.86 mm
Well Diameter - Bottom	6.35 mm
Plate Length	127.8 mm
Plate Width	85.5 mm
Plate Height	14.2 mm
A1 Row Offset	11.2 mm
A1 Column Offset	14.3 mm
Well Center to Well Center Spacing	9 mm
Flange or Skirt Height	6.096 mm
Well Bottom Elevation	3.57 mm
Well Bottom Thickness	0.5 mm
Well Bottom Area	0.3165 cm^2

Kit Components

- 1. <u>Radius[™] 96-well Cell Migration Plate</u> (Part No. 112601): One 96-well black-walled, clear-bottom tissue culture treated plate with each well containing one Radius[™] non-toxic, biocompatible hydrogel spot (96 gel spots total per plate)
- 2. <u>Radius[™] Gel Pretreatment Solution</u> (Part No. 112502): One Sterile Bottle 13.0 mL
- 3. <u>Radius[™] Wash Solution</u> (Part No. 112503): One Sterile Bottle 13.0 mL
- 4. <u>Radius[™] Gel Removal Solution, 100X</u> (Part No. 112504): One Sterile Tube 150 µL
- 5. <u>DAPI Fluorescence Stain, 1000X</u> (Part No. 112002): One Amber Tube $-30 \,\mu L$
- 6. Fixation Solution (Part No. 122402): One Bottle 20.0 mL
- 7. <u>Cell Stain Solution</u> (Part No. 112505): One Bottle 12.0 mL

Materials Not Supplied

1. Adherent migratory cell lines and culture medium



- 2. Multichannel pipette
- 3. Cell culture incubator (37°C, 5% CO₂ atmosphere)
- 4. Inverted light microscope with a digital camera
- 5. Optional: Microscope stage or cage incubator
- 6. Optional: Inverted fluorescence microscope with DAPI filter (350nm/470nm)
- 7. Optional: Imaging Software for measuring cell migration
- 8. Optional: HCS/HCI Instrument

Storage

Upon receipt, aliquot and store the Radius[™] Gel Removal Solution and DAPI Fluorescence Stain at -20°C (avoid multiple freeze/thaw cycles), and transfer the Fixation Solution to 4°C. All other kit components should be stored at room temperature.

Preparation of Reagents

- 1X Radius[™] Gel Removal Solution: Just prior to use, prepare a 1X Radius[™] Gel Removal Solution by diluting the provided 100X stock 1:100 in complete culture medium.
- 1X DAPI Fluorescence Stain: Just prior to use, prepare a 1X DAPI Fluorescence Stain by diluting the provided 1000X stock 1:1000 in PBS.

Assay Protocol (Must be under sterile conditions)

I. Pretreatment of RadiusTM Migration Plate

- 1. Under sterile conditions, remove the Radius[™] 96-well Cell Migration Plate from its packaging. *Note: If wells of the plate have already been used for a previous experiment, the unused wells must be completely moisture-free before proceeding. Please allow the covered plate to dry at room temperature for 1 hour. Each plate can handle up to 3 migration experiment cycles.*
- Determine which wells will be assayed (it is recommended that all samples be tested in triplicate). Slowly add 100 µL of Radius[™] Gel Pretreatment Solution to each well by carefully pipetting down the wall of the well.
 Note: The Radius[™] Gel Pretreatment Solution should <u>not</u> be added to any wells that will not be used immediately.
- 3. Cover the plate and incubate at room temperature for 20 minutes.
- Carefully aspirate the Radius[™] Gel Pretreatment Solution from the wells. Do not allow wells to dry.
 Note: Avoid potential damage to the Radius[™] Gel Spot (located in the center) by aspirating from the edge of the well.
- Slowly add 100 µL of Radius[™] Wash Solution to each well. Proceed to the Cell Seeding Section below.
 Note: Wells can remain in the Radius[™] Wash Solution for up to 1 hour.



II. Cell Seeding

- Harvest and resuspend cells in culture medium at 0.1 0.3 x 10⁶ cells/ml. Note: Cell seeding density is highly cell line dependant, factoring in cell size, spreading and division. Ideally, the desired monolayer confluency at the start of migration (after Radius[™] Gel Removal step) should be 80-90%. Optimization is recommended and can be done in a standard 96-well cell culture plate prior to migration assays.
- Carefully aspirate the Radius[™] Wash Solution from the wells (step 5 above). Do not allow wells to dry.
 Note: Avoid potential damage to the Radius[™] Gel Spot (located in the center) by aspirating from the edge of the well.
- 3. Slowly add 100 μ L of the cell suspension to each well by carefully pipetting down the wall of the well.
- 4. Transfer the plate to a cell culture incubator for 4-24 hours to allow firm attachment/spreading. Take care to avoid shaking or bumping the plate.

III. RadiusTM Gel Removal

- 1. Carefully remove the Radius[™] Migration Plate from cell culture incubator.
- Aspirate the media from each well and wash 3 times with 100 µL of fresh media. Do not allow wells to dry. Note: These washes are intended to remove debris or any dead/unattached cells.
- 3. Prepare sufficient 1X Radius[™] Gel Removal Solution for all wells by diluting the stock 1:100 in culture medium (See Preparation of Reagents Section).
- 4. Aspirate the media from the wells and add 100 μ L of 1X RadiusTM Gel Removal Solution.
- 5. Transfer the plate to a cell culture incubator for 30 minutes to allow complete gel removal.
- 6. Aspirate the 1X RadiusTM Gel Removal Solution from each well and wash 3 times with 100 μ L mL of fresh media. Do not allow wells to dry.
- 7. After the final washing is complete, add 200 μ L of complete medium to each well. Agents that inhibit or stimulate cell migration may also be added directly to the wells.
- 8. At this point, pre-migration images may be captured with an inverted microscope, imaging software, or HCI/HCS instrument.
- 9. Transfer the plate back to the cell culture incubator/microscope stage incubator during the migration process.
- 10. Monitor the migration closure by endpoint or real-time analysis. For time course experiments, live cell compatible dyes or labels are required (e.g. Calcein AM, GFP, RFP). For endpoint experiments, fixed cell detection should be used (Cell Stain, DAPI, TRITC-phalloidin).

IV. (Optional) DAPI Fluorescence Labeling

- 1. Aspirate the media from the wells and add $100 \,\mu$ L mL of Fixation Solution to each.
- 2. Allow the cells to fix for 10 minutes at room temperature. Aspirate and discard the solution.



- 3. Carefully wash each well 3 times with $200 \,\mu\text{L}$ of PBS.
- 4. Prepare sufficient 1X DAPI Fluorescence Stain for all wells by diluting the stock 1:1000 in PBS (See Preparation of Reagents Section).
- 5. Add 100 μ L of 1X DAPI Stain to each well to be stained.
- 6. Incubate 15 minutes at room temperature.
- 7. Carefully wash each well 3 times with $200 \,\mu\text{L}$ of PBS.
- 8. Add 200 µL PBS to each well to keep cells hydrated.
- 9. Examine wells under an inverted fluorescence microscope with DAPI filter (350nm/470nm)

V. (Optional) Cell Staining

- 1. Aspirate the media from the wells and add $100 \,\mu$ L of Cell Stain Solution to each.
- 2. Allow the cells to stain for 15 minutes at room temperature. Aspirate and discard the solution.
- 3. Carefully was each well 3 times with $200 \,\mu\text{L}$ of deionized water.
- 4. Discard all washes and allow wells to dry at room temperature.
- 5. Examine wells under an inverted light microscope.

Analysis of Results

There are a number of software programs available for the analysis of cell migration images. One of these is CellProfilerTM Cell Image Analysis Software offered free-of-charge by the Broad Institute*. You may find more information on this program online at <u>www.cellprofiler.org</u>.

In order to analyze data from our RadiusTM Cell Migration Assays, the CellProfilerTM software must be customized. For your convenience, we have developed add-ons that will customize the program for you. Please visit our website at <u>www.cellbiolabs.com</u>, type "Cellprofiler" in the Search box, and follow the instructions to download the appropriate add-on.

*CellProfiler[™] is a trademark of the Broad Institute. There is no relationship between Cell Biolabs, Inc. and the Broad Institute. Cell Biolabs offers these add-ons as a courtesy to our customers who wish to analyze data obtained using our Radius[™] Cell Migration Assays.



Example of Results

The following figures demonstrate typical results with the RadiusTM 96-well Cell Migration Assay Kit. One should use the data below for reference only. This data should not be used to interpret actual results.



Figure 1. Various Detection Methods with RadiusTM Cell Migration Assay. HeLa cells were seeded overnight. After RadiusTM Gel removal, cells were stained with DAPI, Cell Stain Solution, and Calcein AM (not included in kit) according to the assay protocol.



Figure 2. Cell Migration Time course. HT1080 and NIH3T3 cells were seeded at 20,000 cells/well overnight. After RadiusTM Gel removal, the cells were allowed to migrate for various times.





Figure 3. Inhibition of HT1080 Cell Migration by Cytochalasin D. HT1080 cells were seeded at 20,000 cells/well overnight. After RadiusTM Gel removal, the cells were allowed to migrate for 24 hrs in the presence of various Cytochalasin D concentrations.

References

- 1. Ridley AJ, Schwartz MA, Burridge K, Firtel RA, Ginsberg MH, Borisy G, Parsons JT, Horwitz AR. (2003) *Science* **302**, 1704-9.
- 2. Horwitz R, Webb D. (2003) Curr Biol. 13, R756-9.
- 3. Lauffenburger DA, Horwitz AF. (1996) Cell 84, 359-369.

Recent Citations

- Pearson, P. et al. (2023). Kruppel-family zinc finger proteins as emerging epigenetic biomarkers in head and neck squamous cell carcinoma. *J Otolaryngol Head Neck Surg.* 52(1):41. doi: 10.1186/s40463-023-00640-x.
- Shin, M.D. et al. (2022). Multivalent Display of ApoAI Peptides on the Surface of Tobacco Mosaic Virus Nanotubes Improves Cholesterol Efflux. *Bioconjug Chem.* doi: 10.1021/acs.bioconjchem.2c00371.



- 3. Li, L. et al. (2022). Trehalose Protects Keratinocytes against Ultraviolet B Radiation by Activating Autophagy via Regulating TIMP3 and ATG9A. *Oxid Med Cell Longev*. doi: 10.1155/2022/9366494.
- 4. Wolcott, K.M. et al. (2020). CD34 positive cells isolated from traumatized human skeletal muscle require the CD34 protein for multi-potential differentiation. *Cell Signal*. doi: 10.1016/j.cellsig.2020.109711.
- Toyohara, T. et al. (2020). Patient hiPSCs Identify Vascular Smooth Muscle Arylacetamide Deacetylase as Protective against Atherosclerosis. *Cell Stem Cell*. S1934-5909(20)30157-0. doi: 10.1016/j.stem.2020.04.018.
- 6. Ali, R. et al. (2019). PARP1 blockade is synthetically lethal in XRCC1 deficient sporadic epithelial ovarian cancers. *Cancer Lett*. pii: S0304-3835(19)30538-5. doi: 10.1016/j.canlet.2019.10.035.
- 7. Sapp, R.M. et al. (2019). Circulating microRNAs and endothelial cell migration rate are associated with metabolic syndrome and fitness level in postmenopausal African American women. *Physiol Rep.* **7**(14):e14173. doi: 10.14814/phy2.14173.
- 8. Iyer, D. et al. (2018). Coronary artery disease genes SMAD3 and TCF21 promote opposing interactive genetic programs that regulate smooth muscle cell differentiation and disease risk. *PLoS Genet.* **14**(10):e1007681. doi: 10.1371/journal.pgen.1007681.
- Fraineau, S. et al. (2017). Epigenetic Activation of Pro-angiogenic Signaling Pathways in Human Endothelial Progenitors Increases Vasculogenesis. *Stem Cell Reports*. 9(5):1573-1587. doi: 10.1016/j.stemcr.2017.09.009.
- Wei, W.C. et al. (2017). Functional expression of calcium-permeable Canonical Transient Receptor Potential 4-containing channels promotes migration of medulloblastoma cells. *J. Physiol.* doi:10.1113/JP274659.
- 11. Sidthipong, K. et al. (2016). Rational design, synthesis and in vitro evaluation of novel exomethylene butyrolactone salicyloylamide as NF-kB inhibitor. *Bioorg. Med. Chem. Lett.* doi: 10.1016/j.bmcl.2016.12.017.
- 12. Matijevic Glavan, T. et al. (2016). Toll-like receptor 3 stimulation triggers metabolic reprogramming in pharyngeal cancer cell ine through Myc, MAPK and HIF. *Mol. Carcinog.* doi:10.1002/mc.22584.
- 13. Inoue, K. et al. (2016). Expression of hedgehog signals and growth inhibition by itraconazole in endometrial cancer. *Anticancer Res.* **36**:149-153.
- 14. Tsubamoto, H. et al. (2016). Gremlin 2 is repressed in invasive endometrial cancer and inhibits cell growth in vitro. *Anticancer Res.* **36**:199-203.
- 15. Camacho, M. et al. (2015). Prostacyclin-synthase expression in head and neck carcinoma patients and its prognostic value in the response to radiotherapy. *J Pathol.* **235**:125-135.
- 16. Woodard, G. E. et al. (2014). Characterization of discrete subpopulations of progenitor cells in traumatic human extremity wounds. *PLoS One.* **9**:e114318.
- 17. Felthaus, O. et al. (2014). Migration of human dental follicle cells in vitro. *J Periodontal Res.* **49**:205-212.
- Wong, B. et al. (2013). Adrenomedullin enhances invasion of human extravillous cytotrophoblastderived cell lines by regulation of urokinase plasminogen activator expression and S-nitrosylation. *Biol Reprod.* 88:34.
- 19. Ichikawa, A. et al. (2013). CXCL10-CXCR3 enhances the development of neutrophil-mediated fulminant lung injury of viral and nonviral origin. *Am. J. Respir. Crit. Care. Med.* **187**:65-77.



- Coulouarn, C. et al. (2012). Hepatocyte–stellate cell cross-talk in the liver engenders a permissive inflammatory microenvironment that drives progression in hepatocellular carcinoma. *Cancer Res.* 72: 2533-2542.
- 21. Alcolea, S. et al. (2012). Interaction between head and neck squamous cell carcinoma cells and fibroblasts in the biosynthesis of PGE2. *J.Lipid Res.* **53**:630-642.

Warranty

These products are warranted to perform as described in their labeling and in Cell Biolabs literature when used in accordance with their instructions. THERE ARE NO WARRANTIES THAT EXTEND BEYOND THIS EXPRESSED WARRANTY AND CELL BIOLABS DISCLAIMS ANY IMPLIED WARRANTY OF MERCHANTABILITY OR WARRANTY OF FITNESS FOR PARTICULAR PURPOSE. CELL BIOLABS' sole obligation and purchaser's exclusive remedy for breach of this warranty shall be, at the option of CELL BIOLABS, to repair or replace the products. In no event shall CELL BIOLABS be liable for any proximate, incidental or consequential damages in connection with the products.

Contact Information

Cell Biolabs, Inc. 5628 Copley Drive San Diego, CA 92111 Worldwide: +1 858-271-6500 USA Toll-Free: 1-888-CBL-0505 E-mail: <u>tech@cellbiolabs.com</u> <u>www.cellbiolabs.com</u>

©2013-2024: Cell Biolabs, Inc. - All rights reserved. No part of these works may be reproduced in any form without permissions in writing.

