Cellometer®

Calcein-AM/PI Vitality and Viability Kit

Product Number: CSK-0118 Sample Kit: CSK-0118-S (Not available for purchase)





This product is for RESEARCH USE ONLY and is not approved for diagnostic or therapeutic use

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1.0 Introduction

1.1 Assay Description

Calcein AM (Calcein acetoxymethyl ester) is a cell permeable, non-fluorescent compound. Upon crossing the cell membrane, Calcein AM is rapidly hydrolyzed by cellular esterases inside live cells. The hydrolysis cleaves the AM group, converting the non-fluorescent Calcein AM to a strongly green fluorescing Calcein. The more hydrophilic Calcein is trapped inside the cell (1). Cells that do not possess active cytoplasmic esterases are unable to convert Calcein AM to Calcein, and therefore do not fluorescen green. This allows for a quick and easy detection of metabolically-active cells in a sample.



Stains such as propidium iodide (PI), 7-AAD, and ethidium bromide (EB), are membrane exclusion dyes that are frequently used to stain non-viable nucleated cells with compromised membranes. Acridine orange freely diffuses across the cell membrane and stains DNA in all nucleated cells. When AO and PI are combined it is possible to determine % viability for nucleated cells. When Calcein AM is used in conjunction with PI, it is possible to determine % vitality / viability based on the number of metabolically-active (green fluorescent) and non-viable (red fluorescent) cells in a sample.

Since Calcein AM does not require DNA binding, it stains all metabolically-active cells and can be used to measure metabolic activity in non-nucleated cells, such as platelets (4). Calcein-AM is also a good alternative for analysis of adipocytes, as the AO dye has shown some non-specific binding of lipid droplets that does not occur with Calcein AM (5). Because Calcein AM is photostable, shows low cytotoxicity, does not affect cellular functions, and requires cellular esterases for conversion to green-fluorescing Calcein, it is a popular stain for the examination of cell vitality and viability. (1,2,3).

- 1. Braut-Boucher, F. et al. Journal of Immunological Methods. Vol. 178, Issue 41 (1995).
- 2. Luc S. De Clerck. et al. Journal of Immunological Methods. Vol. 172, Issue 1, (1994).
- 3. Parish, CR. Immunology and Cell Biology. Vol. 77 (1999)
- 4. Verheul , HW. et al. *Blood*. Vol. 96 No. 13 (2000)
- 5. Kilroy, G. et al. PLoS One. Vol.4, Issue 9 (2009)

1.2 Materials Included

- Calcein-AM/PI Vitality and Viability Kit Cat. # CSK-0118 (100 tests)
 - (Component A) Cellometer Propidium Iodide Staining Solution
 - (Component B) Cellometer Calcein-AM Staining Solution

1.3 Additional Materials required, but not included

- dH₂0
- Trypsin EDTA (if working with adherent cells)

1.4 Instrument and Software Requirement

- Cellometer Vision or Vision CBA Image Cytometry System
- Cellometer Vision or Vision CBA Software
- Fluorescence Optics Module VB-535-402 and VB-660-502

2.0 Assay Protocol for CSK-0118

Preparation of Adherent Cells for Staining

- Using 1 x Trypsin-Versene (EDTA), trypsinize cells until they have lifted off the plate (approximately 15 minutes).
- 2. Use the Cellometer Sample Adjustment Calculator to determine the sample volume required to obtain a concentration of 2-3 x 10^6 cells/ mL.
- 3. Spin down cells at 1,000 to 2,000 rpm for five minutes.
- 4. Decant the supernatant and re-suspend cells in 1ml of 1 x PBS or culture media in which the cells were grown.

2.2 Preparation of Calcein AM Reagent

1. Pipette 2 μl Calcein-AM (Nexcelom Part# CSK-0118 Component A) into 18 μl of dH₂O. This is now

Calcein-AM Solution A. Mix by pipetting up and down at least 15 times or vortex.

2.3 Staining Procedure for Cultured Cells

1. Add 5 μl of Calcein-AM Solution A and 5 μl of PI Staining Solution (Nexcelom Part# CSK-0118

Component B) to 40 μl of cell sample.

- 2. Gently pipette the sample up and down ten times, then incubate for 20 min at 37°C in the dark.
- 3. After the 20 minute incubation, the sample is ready for analysis. Proceed to step 2.5.

2.4 Staining Procedure for Whole Blood, Cord Blood, and Bone Marrow Clinical Samples.

- 1. Pipette 10 μ l of fresh blood sample into 70 μ l of 1 x PBS
- 2. Pipette 40 μ l of the diluted blood sample into a new eppendorf tube.
- Add 5 μl of Calcein-AM Solution A and 5 μl of PI Staining Solution(Nexcelom Part# CSK-0118 Component B) to 40 μl of cell sample.
- Gently pipette the sample up and down ten times, then incubate for 20 min at 37°C in the dark.
 After the 20 minute incubation, the sample is ready for analysis. Proceed to step 2.5.

2.5 Data Acquisition

NOTE: Please review the auto save set-up to make sure the acquired data is properly saved (See step 16).

- Gently mix the cell sample by pipetting up and down at least ten times, then load 20 μL into the Cellometer counting chamber and insert into the Cellometer instrument.
- 2. Wait 60 seconds for the cells to settle in the chamber
- 3. Type a name for your sample into the Sample ID text box
- 4. Verify that the default dilution factor for the Calcein AM + PI (For cultured cells) assay is 1.25 by clicking on the pencil icon and locating the "Set Dilution Factor for Assay" in the dialog pop-up screen. The default dilution factor for the Calcein AM + PI CS (Clinical Samples) assay is 10. If a

higher sample dilution was performed in step 2.4, or the cell culture sample was pre-diluted in step 2.3, adjust the dilution factor accordingly. Please note that dilution factor on the main screen is rounded up for display purposes and will read as 1.3.





| $\overline{\blacktriangleright}$ Show Cell Size Distribution Button | |
|---|-----------|
| Set Dilution Factor for Assay | 1.250 |
| Show Percent F1,F2 F1/(F1+F2 | 2)*100% 💌 |

5. Select the Calcein AM + PI or Calcein AM + CS Assay from the Assay drop-down menu in the upper left corner of the main Vision CBA software screen. If this assay is not present in your drop-down menu, import the Calcein AM + PI assay files using the instructions in section 5.0.

| rSETUP | |
|---|------|
| Assay | |
| Calcein AM + Pl 🚽 🄗 🛛 | |
| Assay: Calcein AM - Cell Type Description: | - PI |
| Calcein AM 🥢 | |
| Imaging Mode | |
| Fluorescent 1, Fluorescent 2 | |
| | |

- 6. Unless you are testing clinical samples, the Calcein AM + PI assay should be run with the default software settings. If you suspect that the settings may have been changed, review the default software settings in section 4.0. To update the Calcein AM + PI assay to the Calcein AM + PI CS assay for whole blood, cord blood, and bone marrow clinical samples see section 4.3.
- 7. Click Preview Brightfield Image at the bottom left of the main Vision screen.



8. Turn the focus knob and adjust focus for the bright field image. Cells in focus for the Calcein-AM/PI assay will have a bright center and dark outline. There should be a crisp contrast between background and the cell membrane. See Focus Guide below.



0

0

0

6

- 9. Click Stop Preview
- 10. Click the Preview F1 Image button (bottom left of screen) and verify that the fluorescence signal displays as 100% of range.

For the Calcein AM + PI Assay the default exposure time is 75 milliseconds (msec). For the Calcein AM + PI CS Assay the optimal exposure is 200 milliseconds (msec). Optimal exposure time will generate a bright image with well-defined fluorescent spots.

Under-exposure will yield dark images with weak spots, like the 25 msec image below, with insufficient fluorescent signal. Overexposure will yield images that are too bright with fluorescent



spots that are large and sometimes overlapping, as shown in the 200 ms image below. Spots are also less distinct from background.



- Click Preview F2. The F2 (PI) channel should be set to 2000 msec exposure for the Calcein AM + PI assay and 4000 msec for the Calcein AM + PI SC assay.
- 12. Click the Count button at the bottom of the screen



13. When counting is complete, an initial Results Table will appear on the screen. For optimal counting results, the total cell concentration should be between 3 x 10⁵ cells/mL and 1 x 10⁷ cells/mL (see circled value in the report below). Recount a concentrated or diluted cell sample if necessary.

| Assay: Calcein AM + PI F1 Channel: Calcein AM - Green F2 Channel: Propidium Iodide - Red | | Date: 10/25/2012 14:02:18 | Show Size Distribution |
|--|---|---|------------------------|
| Sample ID: 10uM_75msec_3-2 Dilution: 1.30 Results: | | Instrument Serial #: Vision-301-0088 Instrument Optics: X050-Vx-535-402-Vx-660-502 | |
| Count Calcein positive: 1724 PI positive cells: 1560 Total cells: 3284 Vitality/Viability: 52.6% | Concentration 3.17x10^6 cells/mL 2.86x10^6 cells/mL 6.03x10^6 cells/mL | Mean Diameter 10.7 micron 9.3 micron | |

The default report displays:

- Calcein-positive (metabolically-active) cells counted, cell concentration, and mean cell diameter
- PI-positive (non-viable) cells counted, cell concentration, and mean cell diameter
- Total cells counted and total cell concentration
- % Vitality/Viability for the cell sample: Calcein-positive cells / (Calcein-positive cells + PI-positive cells)
- 14. Click the **Close** button at the bottom right corner of the Counting Results table. Select the F1 Image and check Counted at the right-hand side of the screen.



15. Review the counted image to confirm that the Calcein-positive cells are being counted correctly. Individual cells within clumps should be circled in green, indicating that they are being counted individually. Click F2 Image to confirm that the PI stained cells are being counted correctly as well. If cells are not being counted correctly, please contact Nexcelom Technical Support for assistance with optimization of counting parameters (see Section 3.0).





16. To view bright field and fluorescent cell images that have been saved, open the image folder where the data has been saved. Bright field and fluorescent images are captured for the Calcein AM / PI assay and may be saved automatically. Saved **Raw** images may be opened in the Cellometer software for re-analysis. Images

| Raw image file format .png 💌 Counted image file format .png 💌 |
|--|
| Auto save data.txt |
| File: C: \Users\ Name.NEXCELOM\Desktop Set File |

are only saved if "Auto save data.txt" is selected in the save options menu. The options menu is found by selecting options, then "save options" at the top of the main screen.



A screen-capture software may be used to save both uncounted and counted colorized fluorescent images for presentation and publication.

3.0 Technical Support

- 4.0 Nexcelom Technical Support is available from 9am to 5pm EST.
- 5.0 E-mail: <u>support@nexcelom.com</u>
- 6.0 Phone: 978-327-5340

4.0 Software Settings

4.1 Review Calcein AM/PI Counting Options Screen

Click on the Options Page and select Counting Options. If you suspect that default settings may have been changed, verify that all selections on the instrument screen match the default settings below.

| N Cellometer Vision | CBA 5 - Data Analysis Mode |
|---------------------|------------------------------------|
| Cellomet | Counting Options |
| Visio | Save Options |
| VISIC | Take Background Image |
| r SETUP | Take Fluorescent Background |
| Assay | Change Fluorescence Optics Modules |
| Calcein AM + F | Exposure Adjustment |
| | Instrument |
| Cell Type | |
| Calcein AM | |

4.2 Default (Calcein AM / PI) Software Settings

4.2.1 Check Dialog Screen Settings







2. Verify that all selections on the instrument screen match the default settings below.

| Dialog | | | — ×- |
|-----------------|--|---------------------------|-----------------------|
| Assay Name | Calcein AM + PI | | 🗌 Special Cells |
| | Save as New Assay Type 🗌 Lock As | say from future editing | |
| Description | | | |
| Imaging Mode | Fluorescence 1 (F1) & Fluorescence 2 (F2 |) 💌 🖌 Acquire Brightfield | Image ting |
| -F1 Image | | - F2 Image | |
| Cell Type | Calcein AM | Cell Type PI | • |
| Description: | Edit | Description: | Edit |
| Fluorophore | Calcein AM VB-535-402 💌 | Fluorophore PI | VB-660-502 💌 |
| Fluorescent Exp | 75.0 msec Optics Module | Fluorescent Exp 2000 | .0 msec Optics Module |
| 🗌 Use Br f | Exp Factor of 1.0 | Use Br Exp Factor | of 1.0 |
| Remove | FL Pos from BR count 10.0 | Remove FL Pos fro | om BR count 10.0 |
| Show D | ata File Buttons | | |
| Show S | ample Adjustment Button | | |
| 🔽 Show C | ell Size Distribution Button | | |
| 🔽 Set Dilu | tion Factor for Assay 1.250 | | |
| Show P | ercent F1,F2 F1/(F1+F2)*100% | Use Custom Label | |
| Result Ter | mplate: Calcein AM and PI.rlt_tm | | |
| Set D | efault Browse | | |
| Print Temp | plate: Calcein AM and PI.rlt_tm | | |
| Set D | efault Browse | | |
| FCS Layo | ut File: <none selected=""></none> | | |
| Remo | ve Layout Set FCS Layout | | |
| Print | | | Save Cancel |

4.2.2 Check Cell Type Settings

 Click on the pencil icon under Cell Type on the main Vision CBA software screen to check the Calcein AM cell type settings.



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2. Verify that all selections for the bright field (BR) tab on the instrument screen match the default settings below.

| Cell Type | | X |
|------------|------------------------------|--|
| Cell Type | Name Calcein AM | □ Save as New Cell Type □ Lock from future editing |
| Detailed [| Description | |
| | Brightfield (BR) Fluore | escence (FL) |
| | Minimum Cell Diameter 6. | Maximum 0 micron 30.0 micron |
| | Roundness 0.1 | default: 0.10; range: 0 - 1.0; 1.0 for perfect circle |
| | Contrast Enhancement | ю default: 0.40; range: 0 - 0.8; high value for light cells |
| | Decluster Parameters | |
| | 🗌 Do not decluster clum | 1ps |
| | Decluster 0.1 Edge Factor | 5 default: 0.5; range 0 - 1.0; higher value for more edge enhancement |
| | Deduster Th Factor | default: 1.0; range 0 - 1.0; higher value for more sensitivity |
| | Background 1.0 Adjustment | default: 1.0; range 0 - 1.0; lower value to pick up dim cells |
| | Trypan Blue Viability Para | meters |
| | Minimum | Maximum |
| | Dead Cell 3.0 Diameter | micron 50.0 micron |
| | Sensitivity 1.0 | default: 1.0; range 0 - 6.0; higher value to pick up more dead cells |
| | Uniformity 150 |) default: 150; range 100 - 255 higher value for non-uniform dead cells |
| | Very Dim Dead Ce | lls |
| Prin | t | Save Cancel |

3. Click on the Fluorescence (FL) tab.

| Cell Type | | | |
|-----------|---------------|-------|-------------------|
| Cell Typ | e Name | Calce | ein AM |
| Detailed | Description | | |
| | Brightfield (| (BR) | Fluorescence (FL) |

4. Verify that all selections for the Fluorescence (FL) tab on the instrument screen match the default settings below.

| ell Type |
|---|
| Cell Type Name Calcein AM Save as New Cell Type |
| Detailed Description |
| Brightfield (BR) Fluorescence (FL) |
| Description Calcein AM |
| Minimum Maximum Cell Diameter 6.0 micron 30.0 micron |
| ☐ Normalize intensity for cell size |
| Non-uniform cell fluorescence |
| Roundness 0.10 default: 0.10 range: 0 - 1.0; 1.0 for perfect circle |
| Do not count free nuclei 🛛 Advanced BR/F mode |
| Fluorescence Threshold Parameters |
| C Auto Threshold Fluorescent |
| * Count range 0 - 100% of brightest cell Lower values count dimmer cells |
| Manual Threshold Fluorescent |
| 20.0 * Count range 0 - 100% of image max Lower values count dimmer cells |
| Decluster 1.00 default: 0.9; range 0 - 1.0; Th Factor lower value for better decluster |
| |
| |
| |
| |
| Print Save Cancel |

Nexcelom Bioscience LLC. | 360 Merrimack Street, Building 9 | Lawrence, MA 01843 Telephone: 978.327.5340 | Fax: 978.327.5341 | Email: info@nexcelom.com | www.nexcelom.com 5. Click on the pencil icon under Assay on the main Vision CBA screen, then click the Edit button on the right-hand side of the dialog box.

| SETUP- | Dialog | | - | X |
|------------------------------|-----------------------------|--------------------------------------|---|--------------------|
| Assay | Assay Name | Calcein AM + PI | | Special Cells |
| Calcein AM + PI | Description | Save as New Assay Type Loc | Assay from future editing | |
| | Imaging Mode | Fluorescence 1 (F1) & Fluorescence 2 | (F2) Acquire Brightfield Image Multimode FL Counting | |
| Calcein AM | F1 Image | Coloria AM | F2 Image | |
| Imaging Mode | Description: Fluorophore | Calcein AM Edit | Description: | Edit |
| Fluorescent 1, Fluorescent 2 | Fluorescent Exp | 75.0 msec Optics Module | Fluorescent Exp 2000.0 | msec Optics Module |
| | 🗌 Use Br | Exp Factor of 1.0 | Use Br Exp Factor of | 1.0 |
| | Remov | ve FL Pos from BR count 10.0 | Remove FL Pos from BR co | ount 10.0 |

6. Verify the Brightfield (BR) settings for the F2 image (PI).

| - JPC | | |
|---|--|---|
| Cell Type Name | 기 | Save as New Cell Type |
| Detailed Description | | |
| Brightfield (B | R) Fluorescer | nce (FL) |
| Cell Diame Roundnes Contrast Enhancem Decluster F Do not Decluster Edue Fact Decluster Edue Fact | Minimum ter 6.0 s 0.10 ent 0.40 Parameters t decluster clumps or 0.5 Th 1.0 | Maximum micron 30.0 micron default: 0.10; range: 0 - 1.0; 1.0 for perfect circle default: 0.40; range: 0 - 0.8; high value for light cells default: 0.5; range 0 - 1.0; higher value for more edge enhancement default: 1.0; range 0 - 1.0; higher value for more ensitivity |
| Backgroun Adjustmer | nd 1.0 | default: 1.0; range 0 - 1.0; lower value to pick up dim cells |
| -Trypan Blu | e Viability Parameters | s |
| Dead Cell Diameter | Minimum 3.0 | Maximum micron 50.0 micron |
| Sensitivity | 1.0 | default: 1.0; range 0 - 6.0; higher value to pick up more dead cells |
| Uniformity | 150 | default: 150; range 100 - 255 higher value for non-uniform dead cells |
| Ver | y Dim Dead Cells | |
| Print | | Save Cancel |

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7. Verify the fluorescent settings for the F2 image, by selecting the Fluorescence (FL) tab.

| Cell Type | | | |
|---|--|--|--|
| Cell Type Name Pl | Save as New Cell Type Lock from future editing | | |
| Detailed Description | | | |
| Brightfield (BR) Fluorescence (FL) | | | |
| Description Propidium Iodide (PI) Minimum Maxim Cell Diameter 4.0 micron 3 Normalize intensity for cell size | um 30.0 micron | | |
| Non-uniform cell fluorescence | | | |
| Roundness 0.10 default: 0.10 ra 1.0 for perfect | ange: 0 - 1.0; circle | | |
| Do not count free nuclei Advanced BR/F | ⁼ mode | | |
| C Auto Threshold Fluorescent 10.0 * Count range 0 - 100% of brightest cell Lower values count dimmer cells | | | |
| Manual Threshold Fluorescent 10.0 * Count range 0 - 100% Lower values count dimm | o of image max ner cells | | |
| Decluster 1.00 default: 0.9; rang Th Factor lower value for b | ge 0 - 1.0; etter decluster | | |
| | | | |
| Print | Save Cancel | | |

4.3 Software Settings for Whole Blood, Cord Blood, and Bone Marrow Clinical Samples (Calcein AM + PI CS)

1. Click on the pencil icon under Assay on the main Vision CBA screen.



- 2. Adjust the current settings.
 - 2.1 Check the Save as New Assay Type box
 - 2.2 Rename Assay to Calcein AM + PI CS
 - 2.3 Change the Fluorescent Exp for Calcein AM from 75.0 to 200.0
 - 2.4 Change the Fluorescent Exp for PI from 2000.0 to 4000.0
 - 2.5 Change the Dilution Factor for Assay from 1.25 to 10

2.6 Click Save

| Dialog | | | × |
|-----------------|---|--|---------------|
| Assay Name | Calcein AM + PI | say from future editing | Special Cells |
| Description | | | |
| Imaging Mode | Fluorescence 1 (F1) & Fluorescence 2 (F2) |) Acquire Brightfield Image Multimode FL Counting | |
| F1 Image | | F2 Image | |
| Cell Type | Calcein AM 💌 | Cell Type PI | - |
| Description: | Edit | Description: | Edit |
| Fluorophore | Calcein AM VB-535-402 👻 | Fluorophore PI | VB-660-502 💌 |
| Fluorescent Exp | 75.0 msec Optics Module | Fluorescent Exp 2000.0 msec | Optics Module |
| 🗌 Use Br B | Exp Factor of 1.0 | Use Br Exp Factor of | 1.0 |
| Remove | E FL Pos from BR count 10.0 | Remove FL Pos from BR count | 10.0 |
| Show D | ata File Buttons | | |
| 🗌 Show S | ample Adjustment Button | | |
| 🔽 Show C | ell Size Distribution Button | | |
| 🔽 Set Dilu | tion Factor for Assay 1.250 | | |
| Show P | ercent F1,F2 F1/(F1+F2)*100% | Use Custom Label | |
| Result Ter | nplate: Calcein AM and PI.rlt_tm | | |
| Set D | efault Browse | | |
| Print Temp | plate: Calcein AM and PI.rlt_tm | | |
| Set D | efault Browse | | |
| FCS Layou | ut File: <none selected=""></none> | | |
| Remov | ve Layout Set FCS Layout | | |
| Print | | Save | Cancel |

| Dialog | |
|--|---|
| Assay Name Calcein AM + PI CS | sav from future editing |
| Description | |
| Imaging Mode Fluorescence 1 (F1) & Fluorescence 2 (F2) | Acquire Brightfield Image Multimode FL Counting |
| F1 Image Cell Type Calcein AM | F2 Image Cell Type PI |
| Description: Edit Fluorophore Calcein AM VB-535-402 | Description: Edit Fluorophore PI VB-660-502 💌 |
| Fluorescent Exp 200.0 msec Optics Module | Fluorescent Exp 4000.0 msec Optics Module |
| Use Br Exp Factor of 1.0 | Use Br Exp Factor of 1.0 |
| Remove FL Pos from BR count 10.0 | Remove FL Pos from BR count 10.0 |
| Show Data File Buttons | |
| Show Sample Adjustment Button | |
| ✓ Show Cell Size Distribution Button | |
| Set Dilution Factor for Assay 10 | |

- 3. The updated setting should match those shown below.
- 4. You may now select the Calcein AM + PI CS assay from the drop down menu and proceed with data aquisition in section 2.5.



5.0 Importing a New Cell Type, Assay Type, and Templates

- 1. Click the question mark in the right-hand corner to access the help menu.
- Click on the "Go" button in the "Online Resources" section. This will automatically load the Cellometer Vision CBA Online Resources webpage.
- 3. Under the "Assay Files" tab, locate and download the Calcein AM
 - + PI files onto the desktop.

| -Help Contents | |
|--|----|
| - Introduction and Description - Getting Started - Software Features - Operation Reference - Tutorials | Go |
| Online Resources | |
| - Frequently Asked Questions - Free Training Videos - Contact an Applications Specialist | Go |
| Submit a Support Ticket | |
| - Get Help with Instrument Setup - Get Help with Instrument and Software Issues - Submit Cell Images for Application Support | Go |
| Contact Info | |
| Nexcelom Bioscience LLC 360 Merrimack Street Building 9 Lawrence, MA 01843 USA | |
| Phone: +1 (978)-327-5340 Fax: +1 (978)-327-5341 Email: support@nexcelom.com http://www.nexcelom.com | |
| Done | |

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5.1 Import Result and Print templates

- 1. Navigate to the START menu and select Computer.
- Double click on the C: Drive and locate the ProgramData folder (shown at right), then proceed to Step 3. IF the ProgramData folder is not present, it may be hidden. Follow the instructions below to show hidden folders.



2.1 Click on "Organize" (top left of screen) and select "Folder and search options"



2.2 A Folder Options menu will pop up. Select View.

2.3 Under the "Hidden files and folders" file, select "Show hidden files, folders, and drives"

2.4 Click OK.

- 3. Open ProgramData folder, then open the Nexcelom_VisionCBA folder.
- 4. Open the Template folder. Copy the result template (Calcein AM and PI.rlt_tm) and print template (Calcein AM and PI .prn_tm) files from the new folder on the desktop and paste them into the Template folder.

5.2 Import Cell Type

1. Locate and click on the "Assay Type" at the top of the screen. Followed by Cell Type Manager.



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 Once the Cell Type Manager appears, click on Browse, navigate to the desktop and locate the "Vision_Calcein AM.CellLib" file that has been downloaded onto your desktop.

New folder



3. Select Vision_Calcein AM.CellLib on the desktop, and click Import Highlighted.

| Cell Type Manager | - | _ | × |
|----------------------------|-----------------|----------------------|----------|
| Import from Library | | | |
| O Nexcelom Cell Library | | | |
| | | | Browse |
| Cell Types in Cell Library | | Cell Types in Drop-d | own Menu |
| Calcein AM | | 2372 | * |
| | | 2H3 3T3 | |
| | | 786-0 | |
| | Import | A204 | |
| | Highlighted >>> | A2008 A3B5 | |
| | | A498 | |
| | Import | A549 | |
| | All >> | A549 ATCC | |
| | | Acetic Acid + Pl | |

4. Repeat step 1-4 to download "Vision_PI.CellLib"

5.3 Import Assay Type

1. Locate and click on the "Assay Type" at the top of the screen. Followed by Import / Export



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| Assay Type Manager | | | х | | | |
|--|----------------|--------------------------|---|---|----------------|------------|
| _ Import from Library | | | _ | 1 | Organize 🔻 🛛 🛚 | New folder |
| Nexcelom Assay Library | | | _ | | | |
| C | | Browse |) | | ▲ ★ Favorites | |
| Assays in Assay Library | | Assays in Drop-down Menu | | | Deskton | • |
| 01_BR Concentration 02_Trypan Blue Viability | <u> </u> | 01_BR Concentration | * | | | AT. |
| 03_GFP_Transfection Rate 04_RFP_Transfection Rate | | 03_GFP_Transfection Rate | = | | 🗼 Downloads | -0 |
| 05_YFP_Transfection Rate | E Import | 05_YFP_Transfection Rate | | | 🕮 Recent Plac | es |
| 07_PBMC with AO | Highlighted >> | 07_PBMC with AO | | | Action 1 | |
| 08_PI Viability Jurkat | | 08_PI Viability Jurkat | | | | |
| 10 PI Sperm | All >> | 10 PI Sperm | | | | |
| 11_AOPI_LiveDead | | 11_AOPI_LiveDead | | | | |
| 12_AOEB_LiveDead | | 12_AOEB_LiveDead | | | | |
| 15_Annexin V + PI | | 15_Annexin V + PI | | | | |
| 16_GFP_PI Viability | - | 16_GFP_PI Viability | - | | | |
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3. Select "Vision_Calcein AM + PI.AssayLib" on the desktop, and click Import Highlighted.



4. This assay is now available for use. See section 2.3.4 to select and use assay.

4.0 Additional Resources

4.1 Technical Support

Nexcelom Technical Support is available from 9am to 5:00pm EST.
 E-mail: <u>support@nexcelom.com</u>
 Phone: 978-327-5340

4.3 Storage and Handling

- For long term storage, store CSK-0118 Component A between 2°C and 8°C .
- For long term storage, store CSK-0118 Component B between -24°C and -16°C.

4.4 Warranty

This product is for RESEARCH USE ONLY and is not approved for diagnostic or therapeutic use. Product is warranted to meet the specifications outlined in the Certificate of Analysis when stored and used according to the manufacturer's instructions. No other warranty, expressed or implied (such as merchantability, fitness for a particular purpose, or non-infringement) is granted. Warranty is valid until the expiration date stated on the product label. If no expiration is listed, the warranty is valid for 6 months from the date of product receipt. Warranty will be void if product is stored incorrectly, the recommended protocol is not followed, or the product is used for a different application.

5.0 Ordering Information

5.1 How to Reorder

For orders shipping to destinations in the United States:

- When ordering with a Purchase order
 - Fax a copy of your order to 978-327-5341
 - Email a copy of your order to sales@nexcelom.com
- When ordering with a Credit Card
 - Visit <u>www.shop.nexcelom.com</u> and place your order

For orders shipping to destinations outside the United States:

Contact your local distributor or Nexcelom Representative to place your order