

# BAPTA JF™549 AM

Table 1 - Materials Needed	Most Items Available from ION Biosciences		
Name	Volume	Description/Purpose	
BAPTA JF <sup>TM</sup> 549 AM (50 μg vial)	Dry - 1 Vial	Membrane Permeable Calcium Dye Indicator	
DMSO <sup>1</sup>	25 μL	Solvent for Dissolution of Dye	
100X Pluronic F-127 solution <sup>2</sup>	100 μL	Biocompatible Surfactant for Dye Loading	
100X Probenecid solution (optional) <sup>3</sup>	100 μL	Intracellular Dye Retention Aid	
Assay Buffer - We Suggest One of the Following:			
⇒ 1X HEPES-Buffered Hank's Balanced Salt Solution	10 mL	1X - Ready to Use Assay Buffer	
⇒ 10X Brilliant Assay Buffer	1 mL	10X - Concentrated Assay Buffer	
Water	As Needed	Dilution of 10X Assay Buffer (if used)	

#### Description

BAPTA JF<sup>TM</sup>549 is a red fluorescent (Ex/Em = 546/569 nm), intracellular calcium indicator. Designed with a Janelia Fluor® dye backbone, it displays excellent brightness and photostability for measuring fast calcium dynamics in neurons and cardiomyocytes. The red-shifted fluorescence properties help minimize tissue autofluorescence, and also enable multicolor imaging and the use of optogenetic tools for triggering calcium transients.

### **Laboratory Procedures**

The following protocol provides general guidelines for using this dye to measure intracellular calcium. All loading conditions (dye concentration, temperature, and time) should be optimized for your specific assay, application, and instrumentation.

- 1. Allow all reagents to warm to room temperature before proceeding.
- 2. Add 25  $\mu L$  DMSO<sup>1</sup> to the tube containing BAPTA JF<sup>TM</sup>549 AM.
- 3. Vortex until BAPTA JF<sup>TM</sup>549 AM is fully dissolved. Centrifuge briefly to collect all contents at the tube bottom.
- 4. Add the appropriate volume (see **Table 2**, next page) of 1X Assay Buffer to a 15 mL conical tube. If using 10X Assay Buffer, add the appropriate volume of Water (**Table 2**) and 1 mL of 10X Assay Buffer to a 15 mL conical tube.
- 5. Add 100 µL of 100X Pluronic F-127<sup>2</sup> solution to the conical tube from **step 4**.

Procedure Continues on Next Page





#### Laboratory Procedures (continued)

- 6. (Optional) Add 100 μL of 100X Probenecid<sup>3</sup> solution to the conical tube from **step 5**.
- 7. Vortex conical tube from **step 6** briefly to mix.
- 8. Add the entire contents of the BAPTA JF<sup>TM</sup>549 AM in DMSO solution from **step 3** to the conical tube from **step 7** to make the **Dye Loading Solution**<sup>4</sup>.
- 9. Vortex the **Dye Loading Solution**<sup>4</sup> from **step 8** briefly to mix.

Table 2 Dye Loading Solution <sup>4</sup>	1X Assay Buffer		10X Assay Buffer	
Name	Method A	Method B	Method C	Method D
BAPTA JF <sup>TM</sup> 549 AM in DMSO <sup>1</sup> Solution	25 μL	25 μL	25 μL	25 μL
100X Pluronic F-127 <sup>2</sup> solution	100 μL	100 μL	100 μL	100 μL
100X Probenecid <sup>3</sup> solution	100 μL	-	100 μL	-
1X HEPES-Buffered Hanks Balanced Salt Solution	9.8 mL	9.9 mL	-	-
10X Brilliant Assay Buffer	-	-	1 mL	1 mL
Water	-	-	8.8 mL	8.9 mL
Total	10 mL	10 mL	10 mL	10 mL

- 10. Remove the cell culture medium and add the **Dye Loading Solution**<sup>4</sup> from **step 9**. Recommend volumes are: 35 mm dish or 6-well plate, 1.5 mL; 96-well plate, 100 μL; 384-well plate, 20 μL.<sup>5</sup>
- 11. Incubate in a cell culture incubator at 37°C for 30 minutes.
- 12. Conduct a wash<sup>5</sup> step to remove the **Dye Loading Solution<sup>4</sup>** and replace with cell culture medium or assay buffer. Repeat, if necessary, to completely remove extracellular dye.
- 13. Acquire data using a fluorescence microscope equipped with Cy3 or TRITC filters or a fluorescence plate reader using an excitation wavelength of ~545 nm, an emission wavelength of ~575 nm and an acquisition frequency of 1-10 Hz<sup>6</sup>. See **Table 3** below for recommended settings.

Table 3	Recommended Instrument Settings	
Setting	Recommendation	
Read Mode (Plate Readers)	'Bottom' read mode only	
Ex/Em wavelengths <sup>6</sup>	~545 nm/575 nm	
Cutoff wavelength	570 nm	
Filter selection	Cy3, TRITC	
Contact support@ionbiosciences.com for additional recommendations and guidance on optimizing to your application.		



## Instructions

#### Laboratory Procedures - Footnotes

- DMSO is hygroscopic and should be stored tightly closed. Wet solvent causes difficulties with dissolution of the dye.
- Pluronic F-127 is a biocompatible surfactant that aids in dye dissolution, ensuring equitable dye distribution and cellular loading.
- Probenecid is an anion transport inhibitor that improves intracellular dye retention. Although it is not required for all cell types and dyes, it is recommended in most cases to optimize assay performance.
- <sup>4</sup> The Dye Loading Solution should be used within 2 hours of dye addition for best results.
- In some cases, a no wash format works best. If a no wash format is indicated for your application, we recommend doubling the concentration of 100X Pluronic F-127 and 100X Probenecid in your dye loading buffer.
- To prevent bleed-through or spectral overlap, the Ex/Em wavelengths may need to be optimized by broadening the interval between the wavelengths.

#### Additional Information

Dye indicator and buffer reagents can be purchased either directly from our website or by contacting our Sales Department. Custom and bulk sizes are also available. Contact Sales for more information.

Table 4	Available Reagents	Available Sizes		
Name		Size	Catalog #	
BAPTA JF <sup>TM</sup> 549 AM		500 μg x 1 Vial	1049C	
		50 μg x 10 Vials	1049F	
		50 μg x 3 Vials	1049G	
100X Pluronic F-127	Solution	10 mL Bottle	7601A	
100X Probenecid So	lution	10 mL Bottle	7300P-100	
1X HEPES-Buffered I	Hank's Balanced Salt Solution (1X HHBSS)	100 mL Bottle	7001	
10X Brilliant Assay B	Buffer	10 mL Bottle	7010X	

<sup>\*</sup>JF™ is a trademark owned by the Howard Hughes Medical Institute. Janelia Fluor® is a Registered Trademark of the Howard Hughes Medical Institute.