

# Product Information

## CellBrite™ Fix Membrane Stains

### Kit Contents

Component	Trial size (1 vial*)	Set of 5 vials*
CellBrite™ Fix Membrane Dye	Component A 1 vial	Component A 5 vials
Anhydrous DMSO	99953 150 uL	99953 150 uL

\*Each dye vial makes 20 uL of 1000X dye solution after reconstitution in DMSO.

### Storage and Handling

Store at -20°C, desiccate, and protect from light. Product is stable for at least 12 months from date of receipt when stored as recommended. After reconstitution in anhydrous DMSO, the dye solution can be stored for up to one month at -20°C, protected from light and moisture. Anhydrous DMSO can be stored desiccated at room temperature, 4°C, or -20°C.

### Spectral Properties

Catalog number	Name	Abs/Em*
30090-T, 30090	CellBrite™ Fix 488	480/513 nm
30088-T, 30088	CellBrite™ Fix 555	542/571 nm
30089-T, 30089	CellBrite™ Fix 640	638/667 nm

\*See Figure 1.

### Product Description

CellBrite™ Fix Membrane Stains are a new class of membrane dyes that can be used to stain the cell surface in live cells. Membrane dyes like DiO, Dil, Vybrant® membrane dyes, CellMask™, or PKH dyes can be fixed with formaldehyde. But they are not compatible with detergent permeabilization or methanol fixation, because these treatments extract lipophilic dyes from membranes. In contrast, CellBrite™ Fix Membrane Stains are unique in that their surface staining can withstand permeabilization and methanol fixation, allowing plasma membrane staining to be combined with intracellular immunofluorescence. Unlike lectins such as WGA, which bind specific targets that may vary between cell types, CellBrite™ Fix dyes are general membrane stains.

CellBrite™ Fix Membrane Stains are fluorogenic dyes that rapidly accumulate in the plasma membrane, where they react covalently with the cell surface. As a result, surface staining is well-retained after permeabilization or methanol fixation, with only a slight increase in intracellular fluorescence compared to formaldehyde fixation alone. CellBrite™ Fix dyes have better water solubility than classic lipophilic dyes, and as a result they yield much more uniform staining compared to lipophilic carbocyanine dyes like DiO and Dil. CellBrite™ Fix dyes are non-toxic and do not readily transfer between cells. The dyes have been validated for staining of isolated exosomes for analysis by flow cytometry. They also can be used to stain yeast and bacteria (gram-positive or gram-negative).

CellBrite™ Fix 488 has green fluorescence, CellBrite™ Fix 555 has visible red fluorescence, and CellBrite™ Fix 640 has far-red fluorescence (see Figure 1).

### Considerations for Staining with CellBrite™ Fix Stains

The following are general considerations for using CellBrite™ Fix Stains. See Staining Protocols for step-by-step instructions for use.

- CellBrite™ Fix stains must be used on live cells. The dyes will stain intracellular structures in fixed cells.
- CellBrite™ Fix stains react with proteins and amino acids, therefore, staining must be done in protein- and amine-free buffer such as PBS or HBSS. For adherent cells, we typically use HBSS with calcium/magnesium to maintain cell adhesion and morphology.
- CellBrite™ Fix stains will react with plates coated with poly-L-lysine, collagen, gelatin, or other proteins, resulting in high background. The dyes tend to have higher background on uncoated cell culture surfaces as well. Imaging cells by confocal microscopy can reduce interference from out-of-plane background fluorescence. See tips for imaging, below.
- CellBrite™ Fix dyes react irreversibly with cellular proteins. In live cells, this occurs on the cell surface, because the dyes can't penetrate the membrane. But they do get inside dead cells, where there are many more targets for reaction. As a consequence, the dyes stain dead cells much more brightly than live cells. See tips for imaging, below.
- Cells can be stained in suspension at  $10^5$ - $10^6$  cells in 100 uL following the protocol provided. Pellet the cells by centrifugation and remove the supernatant in between each change of solution.
- CellBrite™ Fix dyes are designed to be fixed shortly after staining, when they primarily localize to the plasma membrane/cell surface. Cells also can be returned to growth medium and cultured after staining, however, dye localization in live cells changes over time. Labeled membranes become internalized, so staining gradually changes from cell surface to intracellular vesicles, usually becoming mostly intracellular after about 24 hours. Internalized CellBrite™ Fix dye is usually detectable for up to 48 hours after staining, though this may vary by cell type.
- Covalent modification of cell surface protein epitopes may interfere with subsequent antibody binding. To reduce the chance of interference, the dye concentration used for labeling should be optimized to use the lowest effective concentration. We also offer MemBrite™ Fix (see Related Products), which are covalent cell surface stains that react with proteins by a different chemistry than CellBrite™ Fix. In cases where CellBrite™ Fix staining interferes with immunostaining for a particular epitope, MemBrite™ Fix may be a suitable alternative.
- See Related Products and visit our website to see our full selection of membrane and cell surface stains, including additional covalent surface stains with more color options, membrane dyes for fixed cells, dyes for long-term membrane staining in live cells, and membrane stains for super-resolution imaging.

### Tips for imaging CellBrite™ Fix staining

#### Confocal vs. epifluorescence microscopy

If you have access to a confocal microscope, we recommend using it to image membrane staining for the best results. Confocal imaging screens out fluorescence from above and below the plane of focus, allowing very crisp imaging of cell boundaries. Compared to regular epifluorescence imaging, confocal is more sensitive and gives you more control over excitation power to limit photobleaching. Membrane dyes can be imaged with a regular epifluorescence microscope, but the images will be more diffuse because fluorescence from membranes above and below the cell borders will be captured.

#### Staining of dead cells

When imaging CellBrite™ Fix staining, do not focus on very bright, rounded-up, or shrunken dead cells. Instead, adjust the plane of focus and imaging settings to detect the live cell membrane staining. The dead cell signal will likely be saturated under these settings. If the dead cell staining interferes with your imaging, try using high magnification and confocal imaging to exclude dead cells from the field of view. Or, try using one of our original CellBrite™ Cytoplasmic Membrane Stains, which do not show dramatic differences in signal between live and dead cells.

## Staining Protocols

### Dye reconstitution

Remove one vial of dye and the anhydrous DMSO from the freezer and bring to room temperature. To make 1000X dye stock solution, add 20  $\mu$ L of anhydrous DMSO to the vial and vortex or pipet up and down to ensure that all of the dye has dissolved. Once dissolved, the dye should be used within a few hours. Unused dye stock solution can be aliquoted and stored desiccated at  $-20^{\circ}\text{C}$  for at least 1 month.

### Mammalian cell staining

1. Wash cells with protein- and amine-free buffer such as PBS or HBSS.
2. Prepare staining solution by diluting CellBrite™ Fix Membrane Dye in buffer to a final concentration of 1X. For example, add 1  $\mu$ L of 1000X dye to 1 mL of buffer. Staining solution should be prepared fresh immediately before use.

**Note:** Dye concentration may need to be optimized for brightness and surface selectivity.

3. Add staining solution to cells and incubate at  $37^{\circ}\text{C}$  for 15 minutes.

#### Notes:

1) Performing dye incubation at  $37^{\circ}\text{C}$  results in strong surface staining, with a small amount of intracellular staining due to dye internalization. Staining also can be performed at room temperature or  $4^{\circ}\text{C}$  to inhibit dye internalization, but incubation time may need to be increased to allow the dye to react with the cell surface.

2) Cells can be incubated with dye at  $37^{\circ}\text{C}$  for longer times without obvious toxicity. However, dye will be internalized and intracellular staining will increase over time.

4. If fixation is not required, cells can be imaged immediately without washing.

**Note:** Cells also can be placed in growth medium for continued culture, but staining will be internalized over time (see Considerations for Staining).

5. To fix cells, wash twice with buffer and fix according to your usual protocol. We usually fix with 4% paraformaldehyde in 1X PBS for 20 minutes at room temperature or  $4^{\circ}\text{C}$ . Cells also can be fixed with pre-chilled methanol for 5-10 minutes at  $-20^{\circ}\text{C}$ . Methanol fixation may result in an increase in intracellular fluorescence.
6. To permeabilize cells, rinse twice with PBS, then incubate with PBS containing 0.1% Triton® X-100 for 10 minutes at room temperature. Permeabilization also can be performed at  $4^{\circ}\text{C}$ . Permeabilization may result in an increase in intracellular fluorescence.

### Staining of bacteria and yeast

CellBrite™ Fix dyes can be used to stain yeast or bacteria (gram-positive and gram-negative), but a higher dye concentration may be needed. We recommend following the same general protocol above, but using 10X dye and optimizing the concentration as needed. Bacteria can be stained at room temperature. Yeast can rapidly internalize the dyes, so staining should be done at room temperature or  $4^{\circ}\text{C}$  to limit staining to the cell surface. Dead cells also may show bright intracellular staining.

## Related Products

Catalog number	Product
30021-30023	CellBrite™ Cytoplasmic Membrane Stains
30070, 30077-30079	CellBrite™ NIR Cytoplasmic Membrane Stains
30105-30109	CellBrite™ Steady Membrane Staining Kits
30092-30099	MemBrite™ Fix Cell Surface Staining Kits
30101-30104	MemBrite™ Fix-ST Cell Surface Staining Kits for STORM
40083	NucSpot® 470 Nuclear Stain for dead or fixed cells
40081	NucSpot® Live 488 Nuclear Stain for live or fixed cells
40082	NucSpot® Live 650 Nuclear Stain for live or fixed cells
40060	RedDot™1 Far-Red Nuclear Stain for live cells
40061	RedDot™2 Far-Red Nuclear Stain for dead or fixed cells
40046	Hoechst 33342, 10 mg/mL in water
70065	LipidSpot™ 488 Lipid Droplet Stain
70069	LipidSpot™ 610 Lipid Droplet Stain
22023	Paraformaldehyde, 4% in PBS, Ready-to-Use Fixative
22020	10X Phosphate-Buffered Saline (PBS)
23001	EverBrite™ Mounting Medium
23002	EverBrite™ Mounting Medium with DAPI
23003	EverBrite™ Hardset Mounting Medium
23004	EverBrite™ Hardset Mounting Medium with DAPI
23008	Drop-n-Stain EverBrite™ Mounting Medium
23009	Drop-n-Stain EverBrite™ Mounting Medium with DAPI

Please visit our website at [www.biotium.com](http://www.biotium.com) for information on our life science research products, including fluorescent CF® dye antibody conjugates, Mix-n-Stain™ Antibody Labeling Kits, apoptosis reagents, and other probes and kits for cell biology research.

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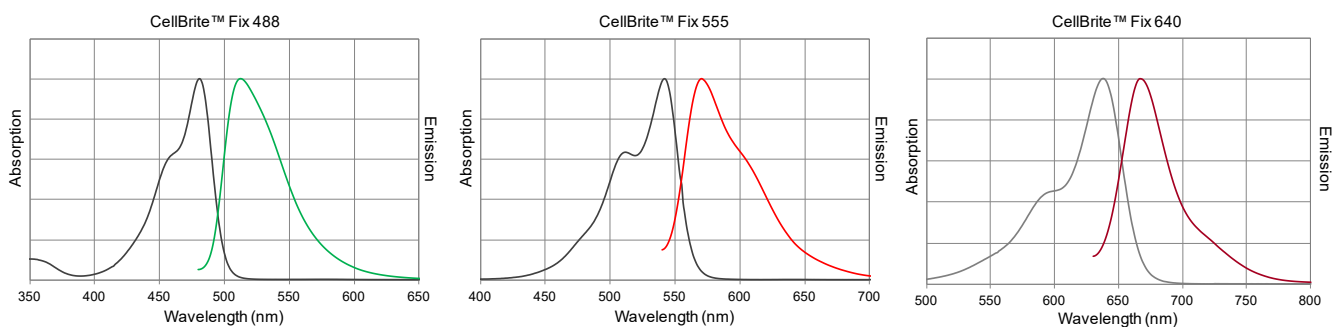


Figure 1. Normalized absorption and emission spectra of CellBrite™ Fix dyes in water.