Product Data Sheet

Tyramide - AcalephFluor633 Reagent (200X)

Catalog #	Source	e Reactivity	Applications
CRG1077		N/A	mIHC
Description		AcalephFluor633 labled Tyra	mide for Multiplex IHC staining or enhanced fluorescent
		IHC staining	
Form		Liquid in PBS	
Directions for	r Use	Add 10 μ l of Tyramide reage	nt into 2 ml of PBS buffer containing 0.003% H2O2. 2 ml
		solution is good for 20 assay	s. Tyramide working solution should be used
		immediately and made fresh	on the day of use.
Platform		Ex/Em = 630/650 nm	
Application		For multiplex immunohistoc	hemical (mIHC) applications, the traditional enzymatic
		amplification procedures are	e sufficient for achieving adequate antigen detection.
		However, several factors lim	it the sensitivity and utility of these procedures.
		Tyramide signal amplificatio	n (TSA) has proven to be a particularly versatile and
		powerful enzyme amplificat	on technique with improved assay sensitivity. TSA is
		based on the ability of HRP,	in the presence of low concentrations of hydrogen
		peroxide, to convert labeled	tyramine-containing substrate into an oxidized, highly
		reactive free radical that car	covalently bind to tyrosine residues at or near the HRP.
		To achieve maximal IHC dete	ection, tyramine is prelabeled with a fluorophore. The
		signal amplification conferre	d by the turnover of multiple tyramide substrates per
		peroxidase label translates u	Itrasensitive detection of low-abundance targets and
		the use of smaller amounts	of antibodies and hybridization probes. In
		immunohistochemical appli	cations, sensitivity enhancements derived from TSA
		method allow primary antib	ody dilutions to be increased to reduce nonspecific
		background signals, and can	overcome weak immunolabeling caused by suboptimal
		fixation procedures or low le	evels of target expression.
Storage/Stab	ility	Store at 4 °C in dark for 1 ye	ar, do not freeze.

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SAMPLE EXPERIMENTAL PROTOCOL

Cell fixation and permeabilization

1. Fix the cells or tissue with 3.7% formaldehyde or paraformaldehyde, in PBS at room temperature for 20 minutes.

- 2. Rinse the cells or tissue with PBS twice.
- 3. Permeabilize the cells with 0.1% Triton X-100 solution for 1-5 minutes at room temperature.
- 4. Rinse the cells or tissue with PBS twice.

Tissue fixation, deparaffinization and rehydration

Deparaffinize and dehydrate the tissue according to the standard IHC protocols. Perform antigen retrieval with preferred specific solution/protocol as needed.

Peroxidase labeling

1. Optional: Quench endogenous peroxidase activity by incubating cell or tissue sample in peroxidase quenching solution (such as 3% hydrogen peroxide) for 10 minutes. Rinse with PBS twice at room temperature.

2. Optional: If using HRP-conjugated streptavidin, it is advisable to block endogenous biotins by biotin blocking buffer.

3. Block with preferred blocking solution (such as PBS with 1% BSA) for 30 minutes at 4°C.

4. Remove blocking solution and add primary antibody diluted in recommended antibody diluent for 60 minutes at room temperature or overnight at 4°C.

5. Wash with PBS three times for 5 minutes each.

6. Apply 100 μ L of secondary antibody-HRP working solution to each sample and incubate for 60 minutes at room temperature.

Note Incubation time and concentration can be varied depending on the signal intensity.

7. Wash with PBS three times for 5 minutes each.

Tyramide labeling

1. Prepare and apply 100 μ l of Tyramide working solution to each sample and incubate for 5-10 minutes at room temperature.

Note If you observe non-specific signal, you can shorten the incubation time with Tyramide. You should optimize the incubation period using positive and negative control samples at various incubation time points. Or you can use lower concentration of Tyramide in the working solution.

2. Rinse with PBS three times.

Counterstain and fluorescence imaging

- 1. Counterstain the cell or tissue samples as needed.
- 2. Mount the coverslip using a mounting medium with anti-fading properties.
- 3. Use the appropriate filter set to visualize the signal from the Tyramide labeling.

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