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Quellenstraße 110, A-1100 Wien T. +43(0)1 489 3961-0 F. +43(0)1 489 3961-7 <u>mail@szabo-scandic.com</u> www.szabo-scandic.com

Cy5.5-SE TEA

®

MedChemExpress

Cat. No.: HY-D0925B	
Molecular Formula: $C_{51}H_{62}N_4O_{16}S_4$	N-0
Molecular Weight: 1115.32	
Target: Fluorescent Dye	
Pathway: Others HO.g ^O	
Storage: Please store the product under the recommended conditions in the Certificate of	Ŋ.
Analysis. 0=\$-	

N-0	

Product Data Sheet

O=S=O OH

BIOLOGICAL ACTIVITY		
Description	Cy5.5-SE TEA (Cyanine5.5 NHS ester TEA) is a CY dye. CY, short for Cyanine, is a compound consisting of two nitrogen atoms connected by an odd number of methyl units. Cyanine compounds have the characteristics of long wavelength, adjustable absorption and emission, high extinction coefficient, good water solubility and relatively simple synthesis ^[1] . CY dyes are of en used for the labeling of proteins, antibodies and small molecular compounds. For the labeling of protein antibodies, the combination can be completed through a simple mixing reaction. Below, we introduce the labeling method of protein antibody labeling, which has certain reference significance ^[2] . Storage: protect from light.	
In Vitro	Protocol1. Protein Preparetion1) In order to obtain the best labeling effect, please prepare the protein (antibody) concentration as 2 mg/mL.2) The pH value of protein solution shall be 8.5±0.5. If the pH is lower than 8.0, 1 M sodium bicarbonate shall be used for adjustment.3) If the protein concentration is lower than 2 mg/mL, the labeling efficiency will be greatly reduced. In order to obtain the best labeling efficiency, it is recommended that the final protein concentration range is 2-10 mg/mL.4) The protein must be in the buffer without primary amine (such as Tris or glycine) and ammonium ion, otherwise the labeling efficiency will be affected.2.Dye Preparation (Cy5.5-SE)Add anhydrous DMSO into the vial of Cy5.5-SE to make a 10 mM stock solution. Mix well by pipetting or vortex.3.Calculation of dye dosageThe amount of Cy5.5-SE required for reaction depends on the amount of protein to be labeled, and the optimal molar ratio of Cy5.5-SE to protein is about 10.Example: assuming the required marker protein is 500 µL 2 mg/mL lgG (MW=150,000), use 100 µL DMSO dissolve 1 mg Cy5.5-SE, the required Cy5.5-SE volume is 5.05 µL, and the detailed calculation process is as follows:1) mol (lgG) = mg/mL (lgG) / MW (lgG) = 2 mg/mL × 0.5 mL / 150,000 mg/monl= 6.7×10-6 mmol2) mol (Cy5.5-SE) = mmol (Cy5.5-SE) / mg/µL (Cy5.5-SE) = 6.7 ×10-6 mmol ×753.88 mg/mmol / 0.01 mg/µL = 5.05 µL (Cy5.5-SE) = mmol (LCy5.5-SE) / mg/µL (Cy5.5-SE) = 6.7 ×10-5 mmol *753.88 mg/mmol / 0.01 mg/µL = 1.05 out (or quere protein samples from denaturation and inactivation.1) A good volume of freshly prepared 10 mg/mL Cy5.5-SE is slowly added to 0.5 mL protein sample Insolution, gently shake to mix, then centrifuge briefly to collect the sample at the bottom o	

	efficiency. 5.Purify the conjugation The following protocol is an example of dye-protein conjugate purification by using a SepHadex G-25 column. 1) Prepare SepHadex G-25 column according to the manufacture instruction. 2) Load the reaction mixture (From "Run conjugation reaction") to the top of the SepHadex G-25 column. 3) Add PBS (pH 7.2-7.4) as soon as the sample runs just below the top resin surface. 4) Add more PBS (pH 7.2-7.4) to the desired sample to complete the column purification. Combine the fractions that contain the desired dye-protein conjugate. MCE has not independently confirmed the accuracy of these methods. They are for reference only.
In Vivo	Cy5.5-labeled factor VIIa is developed for imagining cancer. Cy5.5 labeled with these targeting proteins specifically localize to the tumor xenografts for at least 14 days but unconjugated Cy5.5 does not localize to any xenografts or organs. This method of imaging anti-tissue factor in the tumor VECs may be useful in detecting primary tumors and metastases as well as monitoring in vivo therapeutic responses ^[1] . pH/temperature sensitive magnetic nanogels conjugated with Cy5.5-labeled lactoferrin (Cy5.5-Lf-MPNA nanogels) are developed as a promising contrast agent for preoperative MRI and intraoperative fluorescence imaging of glioma ^[2] . MCE has not independently confirmed the accuracy of these methods. They are for reference only.

CUSTOMER VALIDATION

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Caution: Product has not been fully validated for medical applications. For research use only.

Tel: 609-228-6898 Fax: 609-228-5909

5909 E-mail: tech@MedChemExpress.com

Address: 1 Deer Park Dr, Suite Q, Monmouth Junction, NJ 08852, USA