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Product Data Sheet

BTBCT

Cat. No.: HY-D0038 CAS No.: 525560-81-0 Molecular Formula: $C_{26}H_{15}ClF_6O_6S$

Molecular Weight: 604.9

Target: Fluorescent Dye

Pathway: Others

Storage: Please store the product under the recommended conditions in the Certificate of

Analysis.

BIOLOGICAL ACTIVITY

DescriptionBTBCT is mainly used as a label in time-resolved fluorescence immunoassays (TRFIA). The lower limit of detection for TSH

TR-IFMA is 0.011 mIU/L in a 10 μ l sample volume. The high fluorescence intensity and stability of BTBCT improves the

sensitivity of the assay[1].

In Vitro Protein labeling with BTBCT

1\(\text{Dissolve BTBCT}\) at a concentration of 10 mg/mL in dry ethanol.

2MDissolve the target protein (e.g., BSA or streptavidin) in 0.1 M sodium carbonate buffer to the desired concentration, typically 1 mg/mL.

3 MGradually add the BTBCT solution to the protein solution while stirring.

4\(\text{Maintain stirring at room temperature for 2 hours.}\)

 $5 \boxtimes Filter$ the mixture using an appropriate size filter (usually 0.2 $\mu m).$

6:If necessary, further purify the labeled protein via affinity chromatography to ensure that only functionalized protein is collected.

7:Dialyze the purified protein against a suitable storage buffer (commonly containing 0.05% NaN3).

8:Store at 4°C or freeze as needed.

Indirect Serum TSH TR-IFMA Procedure:

1: Coating the Microplate: Add 30 μ L of coating buffer containing 15 μ g/mL of anti-TSH McAb-05 to each well. Incubate at room temperature for 24 hours. Wash twice with wash buffer.

2: Blocking: Add $40 \mu L$ of blocking buffer (typically containing BSA or another protein) and incubate at room temperature for 6 hours. Wash and air dry.

3:Adding Samples and Standards: Add 10 μ L of TSH standard or serum sample to be tested to each well. Add 10 μ L of a mixture containing biotinylated anti-TSH McAb-04 and McAb-03 at a concentration of about 30 ng/ μ L. Incubate with gentle shaking at room temperature for 1 hour.

 $\hbox{4:Washing: } Thoroughly \ wash \ the \ microplate \ with \ wash \ buffer.$

Adding Signal Generation Reagent: Add 20 μ L of TSH assay buffer containing streptavidin-BSA-BTBCT-Eu complex to each well. Gently shake and incubate for another 20 minutes.

5: Final Washing: Wash four times and rinse twice with distilled water.

6: Fluorescence Measurement: Measure the fluorescence intensity of each well using a time-resolved fluorometer.

Direct Serum T4 TRFIA Procedure:1: Microplate Preparation: Use a microplate coated with anti-T4 antibody, typically at a concentration of $10 \mu g/mL$. Incubate the coated antibody at 4°C overnight.

2: Blocking Non-Specific Sites: Block the microplate with blocking buffer (usually containing 1% BSA) at room temperature for 1-2 hours.

- 3: Adding Samples and Label: Add a predetermined amount of the labeled T4-BSA-BTBCT-Eu complex along with the serum sample to be tested or T4 standard to each well.
- 4: Competition Reaction: Incubate the plate at room temperature for 1-2 hours.
- 5: Washing: Thoroughly wash the microplate with wash buffer.
- 6: Fluorescence Measurement: Measure the fluorescence signal of each well using a time-resolved fluorometer.

MCE has not independently confirmed the accuracy of these methods. They are for reference only.

REFERENCES

[1]. Wu FB et al. new europium beta-diketone chelate for ultrasensitive time-resolved fluorescence immunoassays. Anal Biochem. 2002 Dec 1;311(1):57-67

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

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