

# ReadiLink™ Rapid mFluor™ Red 700 Antibody Labeling Kit \*Microscale Optimized for Labeling 50 µg Antibody Per Reaction\*

Catalog number: 1130 Unit size: 2 Labelings

| Component  | Storage                                    | Amount (Cat No. 1130)                   |
|--|--|---|
| Component A: mFluor™ Red 700                     | Freeze (< -15 °C), Minimize light exposure | 2 vials (One vial is for 50 μg protein) |
| Component B: Reaction Buffer                     | Freeze (< -15 °C), Minimize light exposure | 1 vial (20 μL)                          |
| Component C: TQ <sup>™</sup> -Dyed Quench Buffer | Freeze (< -15 °C), Minimize light exposure | 1 vial (20 μL)                          |

#### **OVERVIEW**

AAT Bioquest's mFluor™ dyes are developed for flow cytometryfocused applications. These dyes have large Stokes Shifts, and can be well excited by the laser lines of flow cytometers (e.g., 405 nm, 488 nm and 633 nm). mFluor™ Red 700 dyes have fluorescence excitation and emission maxima of  $\sim\!657$  nm and  $\sim\!700$  nm respectively. Like Alexa Fluor 700-APC tandem, it has strong absorption at 633 nm, making its conjugates readily excitable by 633 nm red laser. These spectral characteristics make mFluor™ Red 700 an excellent alternative to Alexa Fluor 700-APC tandem. mFluor™ Red 700 is a small organic molecule that is much easier to use than the Alexa Fluor 700-APC tandem. mFluor™ Red 700 SE is reasonably stable and shows good reactivity and selectivity with protein amino groups. This kit has all the essential components to perform two separate labeling reactions with no column purification needed. Each of the two vials of mFluor™ Red 700 provided in the kit is optimized for labeling ~50 µg antibody. mFluor™ Red 700 antibody labeling kit provides a convenient method to label small amounts of monoclonal, polyclonal antibodies or other proteins (>10 kDa) with the mFluor™ Red 700 dye.

### AT A GLANCE

#### **Important**

Warm all the components and centrifuge the vials briefly before opening them. Immediately prepare the necessary solutions before starting your conjugation. The following protocol is a recommendation.

## PREPARATION OF WORKING SOLUTION

## Protein working solution (Solution A)

For labeling 50  $\mu g$  of protein (assuming the target protein concentration is 1 mg/mL), mix 5  $\mu L$  (10% of the total reaction volume) of Reaction Buffer (Component B) with 50  $\mu L$  of the target protein solution.

**Note:** If you have a different protein concentration, adjust the protein volume accordingly to make  $\sim 50~\mu g$  of protein available for your labeling reaction.

**Note:** For labeling 100  $\mu g$  of protein (assuming the target protein concentration is 1 mg/mL), mix 10  $\mu L$  (10% of the total reaction volume) of Reaction Buffer (Component B) with 100  $\mu L$  of the target protein solution.

**Note:** The protein should be dissolved in 1X phosphate buffered saline (PBS), pH 7.2 - 7.4; if the protein is dissolved in glycine buffer, it must be dialyzed against 1X PBS, pH 7.2 - 7.4, or use Amicon Ultra-0.5, Ultracel-10 Membrane, 10 kDa (cat# UFC501008 from Millipore) to remove free amines or ammonium salts (such as ammonium sulfate and ammonium acetate) that are widely used for protein precipitation.

**Note:** Impure antibodies or antibodies stabilized with bovine serum albumin (BSA) or gelatin will not be labeled well.

**Note:** A final protein concentration range of 1 - 2 mg/mL is recommended for optimal labeling efficiency, with a significantly reduced conjugation efficiency at less than 1 mg/mL.

#### SAMPLE EXPERIMENTAL PROTOCOL

#### Run conjugation reaction

 Add the protein working solution (Solution A) to ONE vial of labeling dye (Component A), and mix them well by repeatedly pipetting for a few times or vortex the vial for a few seconds.

**Note:** If labeling 100  $\mu$ g of protein, use both vials (Component A) of labeling dye by dividing the 100  $\mu$ g of protein into 2 x 50  $\mu$ g of protein and reacting each 50  $\mu$ g of protein with one vial of labeling dye. Then combine both vials for the next step.

2. Keep the conjugation reaction mixture at room temperature for 30 - 60 minutes.

**Note:** The conjugation reaction mixture can be rotated or shaken for a longer time if desired.

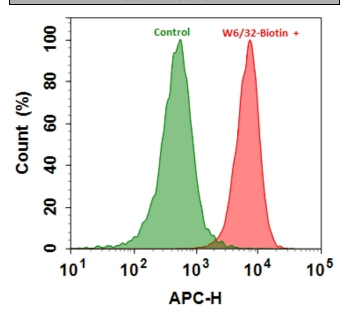
### **Stop Conjugation reaction**

- 1. Add 5 μL (for 50 μg protein) or 10 μL (for 100 μg protein) which is 10% of the total reaction volume of TQ™-Dyed Quench Buffer (Component C) into the conjugation reaction mixture; mix well.
- 2. Incubate at room temperature for 10 minutes. The labeled protein (antibody) is now ready to use.

## Storage of Protein Conjugate

The protein conjugate should be stored at > 0.5 mg/mL in the presence of a carrier protein (e.g., 0.1% bovine serum albumin). For longer storage, the protein conjugates could be lyophilized or divided into single-used aliquots and stored at  $\leq -20$ °C.

### **EXAMPLE DATA ANALYSIS AND FIGURES**



**Figure 1.** HL-60 cells were stained with (Red) or without (Green) 1 μg/mL anti-human HLA-ABC-Biotin (W6/32 mAb). Cells were then incubated with streptavidin labeled using the ReadiLink™ Rapid mFluor™ Red 700 Antibody Labeling Kit (Cat No. 1130). The fluorescence signal was monitored using ACEA NovoCyte flow cytometer in the APC channel.

### **DISCLAIMER**

AAT Bioquest provides high-quality reagents and materials for research use only. For proper handling of potentially hazardous chemicals, please consult the Safety Data Sheet (SDS) provided for the product. Chemical analysis and/or reverse engineering of any kit or its components is strictly prohibited without written permission from AAT Bioquest. Please call 408-733-1055 or email info@aatbio.com if you have any questions.