

### **Product Information & Manual**

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# HyLink<sup>™</sup> Aster 488 Labeling Kit, 100 μg\*10 (SpinDesalt Column)

Cat no. LDG0014RC

# Product Overview Package component

Package	(100 μg x 10)	Storage
Aster 488	10 vials	-20°C
10X Modifier	1 vial	-20°C
10X Quencher	1 vial	-20°C
SpinDesalt		
Column	10 vials	4°C
(LDG0008RC)		

#### Description

Leadgene HyLink<sup>TM</sup> Aster 488 Conjugation Kit is designed for small scale conjugation. The NHS ester groups of Aster 488 can react with amino group to form a stable amide group. It provides a rapid and easy process with high efficiency and fluorescent intensity to conjugate antibodies or protein to Aster 488. The total process completed in 3 hours and less than 30 minutes hands-on time.

#### **Procedure**

- (1) Equilibrate reagent to room temperature before using. Make sure all buffers are well dissolved. If not, please vortex the vial to make salts dissolved.
- (2) Dissolve antibody in PBS or other buffer that do not contain amine, tris, NaN3 or glycerol. Add 10X Modifier to antibody (e.g. 1 μL of 10X Modifier for 9 μL of antibody).
- (3) Spin down the vial of Aster 488 before using.
- (4) Open the cap of the vial of Aster 488 and pipette antibody into the vial. Mix gently by pipetting several times until Aster 488 dye is well dissolved.

- (5) Cover the cap on the vial and spin down the vial. Incubate in the dark at room temperature for 2 hours.
- (6) Method 1:

Using SpinDesalt Column to remove unconjugated Aster 488. Please refer to the protocol of SpinDesalt Column, catalogue no. LDG0008RC. Collect labeled antibody and stabilize with 1% bovine serum albumin or another stabilizer.

#### Method 2:

Add 10X Quencher to Ab-Fluorescent mixture. Incubate in the dark at room temperature for 30 minutes. Stabilized with 1% bovine serum albumin or another stabilizer.

- (7) Store the labeled protein protected from light.
- (8) For protein conjugation, it can be calculated by formula below:

Quantities of protein = quantities of kit (e.g. 100  $\mu$ g) x (M.W.of target protein)/(150000 (M.W.of  $\mu$ g)

#### Important notes

(1) Antibody concentrations of 0.5-2 mg/mL generally give optimal results.

Kit size	Antibody amount	Reaction volume
100 μg x 10	50-200 μg	40-200 μL

(2) Common non-buffering salts (e.g. sodium chloride) have no effect on conjugation efficiency. Avoid buffer component that contains primary amine (e.g. amino acid or ethanolamine) and thiols (e.g. mercaptoethanol or DTT).



#### Determine of DOL (degree of labeling)

(1) Calculate concentration of protein:

Portein concentration (M) = 
$$\frac{A_{280}\text{-}(A_{max} \times CF)}{\epsilon_{protein}} \times Dilution factor$$

(The molar extinction coefficient of IgG is 210000 M<sup>-1</sup> cm<sup>-1</sup>.)

Dye	Amax /	CF (Correcti	Extinction
	Emission	on factor)	coefficient (ε) M <sup>-1</sup>
			cm <sup>-1</sup>
Aster 488	490 / 515	0.1	70000

(2) Calculate DOL:

$$DOL = \frac{A_{max} \text{ of labeled protein}}{\epsilon_{Aster \, 488} \times Protein \, concentration \, (M)} \times Dilution \, factor$$

#### Disclaimer

This product is for research use only and is not intended for diagnostic use.

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<sup>\*</sup>  $\epsilon_{protein}$ : protein molar extinction coefficient.



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### SpinDesalt Column

Cat no. LDG0008RC

#### **Product Overview**

#### **Package**

SpinDesalt Column, 0.5 mL, 5 columns

#### Introduction

Leadgene SpinDesalt Column is a convenient and rapid product that is suitable for desalting, buffer exchange, or removal of small molecules from proteins or other macromolecular samples with a molecular weight larger than 5 kDa. The SpinDesalt Column is filled with 0.5 mL of Smartdex G-25, which efficiently separates proteins from small molecules (such as salt and other small molecules like biotin).

#### Storage

SpinDesalt Column should be store at 2-8°C.

#### **Procedure**

#### Preparation of the equilibration buffer

The equilibration buffer can be chosen according to the buffer you want to replace. It's recommended to filter it with a 0.22  $\mu m$  or 0.45  $\mu m$  membrane before using the buffer.

#### Preparation of the sample

It is recommended to centrifuge the sample or filter it with a  $0.22 \, \mu m$  or  $0.45 \, \mu m$  membrane before loading, to reduce the impact of contaminant.

#### SpinDesalt Column protocol

(1) Prepare a SpinDesalt Column by breaking off the bottom closure and placing the column into a 2 mL

collection tube.

- (2) Centrifuge the column at 1,000 × g for 1 minute, discard the storage buffer and return column to the same collection tube.
- (3) Adding 0.25 mL of the equilibration buffer to the top of the resin bed and centrifuging at  $1,000 \times g$  for 1 minute. Discard the flowthrough and repeat this step 3 times.
- (4) Place the column into a new 1.5 mL collection tube and apply approximately 0.1-0.2 mL of the sample directly onto the resin bed. Centrifuge the column at 1,000 × g for 1 minute.
- (5) The collected flowthrough solution is the purified sample.

#### Important notes

- (1) Please break off the bottom closure before using the column.
- (2) Please centrifuge the column at 1,000 x g.
- (3) Each column can process a 100-200  $\mu$ L sample at a time.
- (4) The recovery rate of the SpinDesalt Column is related to the type of protein and other biomolecules, usually exceeding 85%. Increasing the sample concentration or volume can improve the recovery rate.
- (5) If sample < 0.1 mL, please use the equilibration buffer to adjust the volume to at least 0.1 mL to increase the recovery rate.
- (6) The resin bed of SpinDesalt Column can be temporarily stored in the equilibration buffer.



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