R-Phycoerythrin Labeling Kit - NH₂

Technical Manual

"echnical Manual (Japanese version) is available at http://www.dojindo.co.jp/manual/ik23.pc

General Information	R-Phycoerythrin Labeling Kit - NH ₂ is primarily used for the preparation of R-phycoerythrin-labeled antibody for im- munostaining and cellular proteins for tracing. NH ₂ -Reactive R-Phycoerythrin, a component of this kit, has succin- imidyl ester groups, and can easily make a covalent bond with amino groups of IgG or other proteins without any activation process. Filtration Tube included in this kit is used for removing small molecules in the sample protein such as Tris buffer and amine compounds that interfere with the assay or labeling reaction. The maximum excita- tion and emission wavelengths of the R-phycoerythrin-labeled proteins are 564 nm and 575 nm, respectively. This kit contains all of the necessary reagents for labeling, including the storage buffer for conjugate.
Kit Contents	- NH2-Reactive R-Phycoerythrin3 tubes- WS Buffer4 ml x 1- Reaction Buffer200 µl x 1- Filtration Tube3 tubes
Capacity	Three samples labeling - sample requirement : molecular weight > 50,000; amount: 50-200 μg
Storage Condition	Store at 0-5°C. This kit is stable for 1 year at 0-5°C before opening.
	Caution After a NH -Reactive R-Phycoerythrin is taken out from the seal bag, keep the unused NH -Reactive R-Phycoerythrin(s) in the bag, seal tightly and store at -20°C. Store the other components at 0-5°C.
Required Equipment	 - 10 μl and 200 μl adjustable pipettes - Incubator (37°C) - Microtubes
Precaution	 If the target protein solution contains other proteins with molecular weight larger than 10,000, such as serum abumin or gelatin, purify the protein solution, and use the purified target proteins for R-phycoerythrin labeling, because it might interfere the filtering or labeling reaction. If the protein solution contains small insoluble materials, centrifuge the solution, and use the supernatant for the labeling. The droplets which induced from the dry inhibitor of membrane, are occasionally found inside Filtration Tube while storing the kit at 0-5°C or after returning to room temperature. This phenomenon does not affect the performance. This kit includes microtubes containing solutions. Since there is a possibility that the droplets might attach to the the solution endpering the solution provide the droplets might attach to the droplets which induced from the droplets. Since there is a possibility that the droplets might attach to the the effect of the performance. The droplets which induced from the droplets is a possibility that the droplets might attach to the the effect of the performance. The kit includes microtubes containing solutions. Since there is a possibility that the droplets might attach to the the effect of the performance.
	Maximum excitation wavelength : 564 nm Maximum emission wavelength : 575 nm
	Excitation and emission spectra of R-phycoerythrin-labeled protein LK23: R-Phycoerythrin Labeling Kit - NH2

General Protocol

- Labeling for IgG -



Step 1. Add 100 μI WS Buffer and the sample solution containing 50 - 200 μg IgG^{a)} to a Filtration Tube.



Step 2. Pipette to mix and centrifuge at $8,000 \times g$ for 10 min.^{b)}



Step 3. Add 100 µl WS Buffer to a Filtration Tube again.



Step 4. Centrifuge at 8,000 x g for 10 min again.^{b)}



Step 5. Add 10 μI Reaction Buffer to NH_2 -Reactive R-Phycoerythrin, and dissolve with pipetting. $^{\circ)}$



Step 6. Add NH₂-Reactive R-Phycoerythrin solution to the IgG concentrated on the Filtration Tube



Step 7. Incubate the tube at 37°C for 2 hours after pipetting to mix.



Step 8. Add 190 µI WS Buffer, and pipette about 10 times to recover the conjugate.^{d)} Transfer the solution to a microtube (not included in this kit), and store at 0-5°C.^{e)}

- a) The volume of IgG solution should be less than 100 µl. If the IgG concentration is lower than 0.5 mg/ml, repeat Steps 1 and 2 until the total IgG accumulation becomes 50 200 µg.
- b) If solution still remains on the membrane after the centrifugation, spin for another 5 min.
- c) NH_-Reactive R-Phycoerythrin can be hydrolyzed by water. Proceed to Step 6 immediately after the preparation of the NH_-Reactive R-Phycoerythrin solution.
- d) One to two R-phycoerythrin should be introduced into one IgG molecule. Unconjugated R-phycoerythrin remained in the solution might cause background increase with immunoassay. If purification is necessary, purify the conjugate using a gel permeation column or an affinity column for IgG.
- e) We recommend using WS Buffer to recover the conjugate. However, you can use an appropriate buffer for the downstream experiments.

Q & A • Can I use this kit to label antibody which is commercially available?

Yes. However, if antibody solution contains other proteins such as serum albumin or gelatin, labeling reaction might be interfered by that protein. Purification of the antibody solution with affinity chromatography is necessary prior to use this kit. Contact us for the purification procedure, if you need.

- How long is the R-phycoerythrin labeled protein stable?
 - The stability depends on the protein itself. For longer storage, add equal volume of glycerol to the sample solution and store at -20°C.
- What is the minimum amount of protein that can be labeled using this kit?
 We recommend using 50 µg as a minimum amount. Though 10 µg protein can be labeled using this kit, the background might be increased.
- Does NH₂-Reactive R-Phycoerythrin form an oligomer during the labeling reaction?
 No. Since all amino groups of NH₂-Reactive R-Phycoerythrin are blocked, no oligomerization is occurred.
- Can I use the R-Phycoerythrin conjugated protein that is precipitated in storage?
 Yes. The precipitated protein should be removed by centrifugation at 10,000 x g for 10 min, and use the supernatant.
- Is there any notice for treatment of living cells with the R-Phycoerythrin conjugated protein?
 We recommend using PBS including 2-10% FBS for preparation of cell suspension to maintain the best cell condition.
- Does recovery buffer (WS Buffer) have harmful effect to living cells?
 No. WS Buffer contains stabilizing agent (surfactant) that is controlled of its concentration without cytotoxicity.
 If you are concerned about the additive in WS Buffer, you can use your own buffer currently used instead of WS Buffer.

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