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# S9351S

50 nmol Lo	ot: 0021202
Store at: -20°C Ex	cp: 2/14

## Introduction

CoA Biotin is a non cell-permeable substrate based on biotin with an amidocaproyl linker. It is suitable for applications such as biotinylation of ACP-tag or MCP-Tag fusion proteins on the surface of living cells for detection with streptavidin fluorophore conjugates. This package contains 50 nmol of CoA Biotin substrate, sufficient to make 10 ml of a 5  $\mu$ M solution for the labeling of ACP-tag or MCP-tag fusion proteins on cells.

The ACP-tag and MCP-tag are polypeptide tags (8 kDa) based on the acyl carrier protein. MCP-tag contains two mutations (D36T and D39G). Both allow the specific, covalent attachment of virtually any molecule to a protein of interest. Substrates are derivates of coenzyme A (CoA). In the labeling reaction, the substituted phosphopantetheine group of CoA is covalently attached to a conserved serine residue of the ACP-tag or the MCP-tag by a phosphopantetheinyl transferase (SFP Synthase, or ACP Synthase).

While ACP Synthase (NEB #P9301) will preferentially modify the ACP-tag, SFP Synthase (NEB #P9302) will label both ACP-tag and MCP-tag.

Having no cysteines, the ACP-tag and the MCP-tag are particularly suited for specifically labeling cellsurface proteins, and should be useful for labeling secreted proteins with disulphide bridges such as antibodies.

There are two steps to using this system: subcloning and expression of the protein of interest as an ACP-tag or MCP-tag fusion, and labeling of the fusion protein using the appropriate synthase with the CoA substrate of choice. Expression of ACP- and MCP-tagged proteins is described in the documentation supplied with the pACP and pMCP plasmids, respectively. The labeling of the fusion proteins with the CoA Biotin is described below.

# Materials required but not supplied:

ACP Synthase (NEB #P9301) for labeling ACP-tag SFP Synthase (NEB #P9302) for labeling ACP-tag or MCP-tag

Cells expressing ACP-tag or MCP-tag fusion proteins

Tissue culture materials and media

Transfection reagents

Fluorescence microscope with suitable filter set DMSO

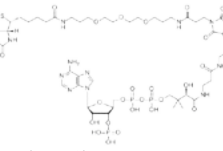


Figure 1. Structure of CoA-Biotin (MW 1365.3 g/mol)

# Storage

CoA Biotin should be stored at  $-20^{\circ}$ C (long term) or at 4°C in the dark (short term, less than 4 weeks). Protect the substrate from light and moisture. With proper storage at  $-20^{\circ}$ C the substrate should be stable for at least one year dry or 3 months dissolved in DMSO.

## **Quality Controls**

Purity and Characterization: Purity of CoA-Biotin was determined to be 94% by HPLC analysis. Molecular weight [M]<sup>-</sup> was determined by MS to be 1363.4 (1363.4 expected).

*In vitro* protein labeling: Reaction of CoA-Biotin (10  $\mu$ M) with purified ACP-MBP (Maltose Binding Protein, 5  $\mu$ M) and ACP Synthase (1  $\mu$ M) *in vitro*, followed by mass spec analysis, indicated an efficiency of labeling of  $\geq$  95%.

## Instructions for Cellular Labeling

ACP-tag and MCP-tag fusion proteins are expressed by transient transfection. For expression of fusion proteins with the ACP-tag or MCP-tag refer to instructions supplied with the pACP and pMCP plasmids. For cell culture and transfection methods, refer to established protocols.

Dissolve one vial of CoA substrate (50 nmol) in 50  $\mu$ l of DMSO to give a labeling stock solution of 1 mM CoA substrate. Mix for 10 minutes until all

the substrate is dissolved. Store this stock solution in the dark at 4°C, or for extended storage at  $-20^{\circ}$ C. Different stock concentrations can be made, depending on your requirements. The substrate is soluble up to at least 10 mM.

#### Protocol for Labeling reaction:

- 1. Dilute the Substrate Stock Solution 1:200 in medium to a final concentration of 5  $\mu$ M. Mix substrate with medium thoroughly by pipetting up and down 10 times. For best performance, add the CoA substrate to complete medium, including serum (0.5% BSA can be used for experiments carried out in serum-free media). Add MgCl<sub>2</sub> to a final concentration of 10 mM. Finally, add ACP Synthase or SFP Synthase to a final concentration of 1:40. Do not prepare more medium with substrate, MgCl<sub>2</sub>, and synthase than you will consume within one hour.
- 2. Replace the medium on the cells expressing an ACP-tag or MCP-tag fusion protein with the labeling medium and incubate at 37°C, 5%  $CO_2$  for 30 minutes.

Number of wells in plate	Recommended Volume for Cell Labeling
6	1 ml
12	500 µl
24	250 µl
48	100 µl
96	50 µl

These recommendations are for culturing cells in polystyrene plates. For confocal imaging, we recommend using chambered coverglass such as Lab-Tek II Chambered Coverglass which is available in a 1, 2, 4 or 8 well format from Nunc (www.nuncbrand.com).

3. Wash the cells three times with tissue culture medium with serum.

# Microscopy

## <u>Fixation</u>

After labeling the ACP-tag or MCP-tag fusion proteins with CoA-Biotin, the cells can be fixed with 3.3% para-formaldehyde which does not result in the loss of signal. Avoid fixation using

ethanol as this may lead to a high background staining of endogenous biotinylated proteins found preferentially in mitochondria.

## **Detection**

To visualize the ACP-tag or MCP-tag fusion protein *in-situ*, permeabilize the cells with 0.5% Triton in PBS and block the cells with 1% BSA in PBS containing 0.5% Triton. Incubate the fixed cells with an appropriate streptavidin/avidin conjugate (e.g. streptavidin-fluorophore) and image the cells according to the instructions supplied with the conjugate.

# Western Blotting

Biotinylated proteins from cell lysates can be visualized on Western Blots using standard streptavidinbased detection reagents. For Western blotting experiments it may be more efficient to label the ACP-tag or MCP-tag fusion proteins after lysis of the cells.

# **Optimizing Labeling**

Optimal substrate concentrations and reaction times range from  $5-30 \ \mu$ M and 30 minutes to overnight, respectively, depending on experimental conditions and expression levels of the ACP-tag or MCP-tag fusion protein. Best results are usually obtained at concentrations between 5 and 10  $\mu$ M substrate and 30 minutes reaction time. Increasing substrate concentration and reaction time usually results in a higher background and does not necessarily increase the signal to background ratio.

# Stability of Signal

The turnover rates of the ACP-tag or MCP-tag fusion protein in live cells under investigation may vary widely depending on the fusion partner. We have seen half-life values ranging from less than one hour to more than 12 hours. Where protein turnover is rapid, we recommend processing the cells for imaging or blotting immediately after the labeling reaction.

# Counterstaining

Cells can be counterstained with any live-cell dye that is compatible with the properties of the ACP-tag or MCP-tag substrate for simultaneous microscopic detection. We routinely add 5  $\mu$ M Hoechst 33342 (nuclear stain) after labeling with avidin/streptavidin followed by 2 short washing steps. Counterstaining of cells is also possible with dyes that do not enter live cells after fixation and permeabilization.

(see other side)

#### Immunocytochemistry

Antibody labeling of the fusion protein can be performed after ACP-tag or MCP-tag labeling and fixation of the cells according to standard protocols without loss of the ACP-tag or MCP-tag signal.

## Troubleshooting

## No Labeling

If no labeling is seen, there is probably a problem with the expression of your fusion protein. Verify your transfection method to confirm that the cells contain the fusion gene of interest. If this is confirmed, check for expression of the ACP-tag or MCP-tag fusion protein. A pACP or pMCP control plasmid may also be used as a positive control.

# Weak Labeling

Weak labeling may be caused by insufficient exposure of the fusion protein to the substrate. Try increasing the concentration of CoA Biotin substrate and/or the incubation time. Improving the protein expression may also improve the signal. If the protein has limited stability in the cell, it may help to analyze the samples immediately after labeling.

# Instructions for Labeling of Proteins in Solution:

Dissolve the vial of CoA Biotin (50 nmol) in 17  $\mu$ l of DMSO to yield a stock solution of 3 mM substrate. Mix for 10 minutes until all the substrate is dissolved.

# Protocol for Labeling Reaction:

- Prepare a protein solution containing up to 20 μM ACP-tag or MCP-tag fusion protein to be labeled in an appropriate buffer.
- 2. Add 3 mM ACP-tag or MCP-tag substrate solution to a total volume of 1% of the volume of the protein solution. Carefully pipette the material up and down to mix, and vortex briefly.
- 3. Incubate for 1 hour at 25°C. Alternatively incubate overnight at 4°C.

Biotin labeled proteins can be used for Western Blot analysis. Alternatively they can be used to study protein interaction in interaction assays or used in pull-down assays. Biotinylated proteins can also be immobilized on streptavidin beads or on streptavidin plates.

# Removal of Unreacted Substrate (optional)

After the labeling reaction you may wish to separate the unreacted substrate from the biotinylated fusion protein before using it for your experiment. You can use size-exclusion columns or extensive dialysis. Please refer to the vendor's instructions for the separation tools you are using. Removal of unreacted substrate should not be necessary prior to detection of biotin labeled ACP-tag or MCP-tag fusion proteins on Western blots.

# Notes for Labeling in Solution

## Labeling

ACP-tag or MCP-tag fusion proteins can be purified before labeling, but the labeling reaction also works in non-purified protein solutions (including cell lysates).

# **Confirmation of Labeling by Western Blot Analysis**

Labeled ACP-tag or MCP-tag fusion proteins can be easily analyzed on a SDS-PAGE gel/Western blot analysis because the covalently bound label will remain attached to the protein. The biotin label can be detected on an SDS-PAGE gel followed by Western Blot using a Horseradish Peroxidase or Alkaline Phosphatase labeled avidin/streptavidin (e.g. streptavidin-HRP) and the corresponding detection method as described by the supplier of the enyzme conjugate.

## **Troubleshooting for Labeling in Solution**

# Labeling Reaction

If solubility problems occur with your ACP-tag or MCP-tag fusion protein, we recommend testing a range of pH (pH 5.0–pH 10.0) and ionic strengths. The salt concentration may also need to be optimized for your particular fusion protein (50–250 mM).

If stickiness of the fusion protein is a problem we recommend adding Tween 20 at a final concentration of 0.05% to 0.1%. The ACP-tag or MCP-tag activity is not affected by this concentration of Tween 20.

If exhaustive labeling of a protein sample is not achieved using the recommended conditions, try the following protocol modifications: Double the incubation time to two hours total at 25°C or to 24 hours at 4°C; or halve the volume of protein solution labeled (50  $\mu$ l of a solution containing up to 20  $\mu$ M ACP-tag or MCP-tag fusion protein). Both approaches may be combined.

If your fusion protein is particularly sensitive to degradation or to loss of activity, you can try reducing the labeling time or decreasing the labeling temperature. If you label at 4°C we recommend overnight incubation.

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