



A Geno Technology, Inc. (USA) brand name

HOOK™ Sulfo-NHS-LC-Biotin

For the coupling of biotin to protein amine groups

(Cat. # BS-07)



INTRODUCTION
ITEMS SUPPLIED
STORAGE CONDITION
SPECIFICATIONS
PRECAUTIONS
ADDITIONAL ITEM(S) REQUIRED
PREPARATION BEFORE USE
A. PROTEIN SAMPLE PREPARATION
B. CALCULATION OF QUANTITY OF BIOTIN AGENT NEEDED FOR CONJUGATION5
C. PREPARATION OF BIOTIN AGENTS
D. BIOTIN CONJUGATION REACTION
E. REMOVAL OF UNCONJUGATED BIOTIN AGENTS
F. ESTIMATION OF BIOTIN INCORPORATION EFFICIENCY
IMPORTANT INFORMATION
PROTOCOL 1: CUVETTE PROTOCOL
PROTOCOL 2: MICROPLATE PROTOCOL
CALCULATIONS8
TROUBLESHOOTING
APPENDIX 1: SAMPLE EQUILIBRATION WITH TUBE-O-DIALYZER™ (NOT SUPPLIED) 10
APPENDIX 2: INSTRUCTIONS FOR CELL SURFACE PROTEIN BIOTINYLATION11
CELL SAMPLE PREPARATION:
BIOTIN AGENT & CELL SURFACE REACTION
DELATED DECOLICTS

INTRODUCTION

This kit is designed for the conjugation of biotin to protein primary amine groups and is supplied with the biotin reagent, a specific Optimizer Buffer^m, for enhanced conjugation, Spin- OUT^{m} columns, for purification of labelled protein, and reagents to determine the amount of biotinylation.

Biotin, a 244 Dalton molecule, exhibits an extraordinary binding affinity for avidin and streptavidin (K_a =10¹⁵ M⁻¹). The biotinylated molecules are efficiently probed with avidin or streptavidin conjugated to reporter molecules, such as peroxidases or phosphatases. The use of biotin labeled proteins in ELISA, Western blotting and dot blotting is a popular technique.

The most widely used amine reactive biotinylation reagents are the water insoluble N-hydroxysuccinimide (NHS) esters or the water soluble N-hydroxysulfosuccinimide (sulfo-NHS) esters. The addition of a charged sulfonate (SO_3) on the N-hydroxysuccinimide ring of the sulfo-NHS esters results in their solubility in water (~10mM), but not permeable to plasma membranes. The solubility and impermeability to plasma membranes makes them ideal for studying cell surface proteins as they will only react with the protein molecules on the outer surface of plasma membranes. The reactions of the NHS and sulfo-NHS esters with amines are virtually identical leading to the formation of an amide bond and release of NHS or sulfo-NHS.

HOOK™ Sulfo-NHS-LC-Biotin kit is designed for the coupling of 1-10mg protein in 1ml buffer, suitable for 10 couplings.

ITEMS SUPPLIED (Cat. # BS-07)

Description	Size
HOOK™ Sulfo-NHS-LC-Biotin Agent	25mg
Optimizer Buffer™ I [5X]	2 x 25ml
Spin- <i>OUT</i> [™] GT-600, 5ml	10 columns
OneQuant™ HABA/Avidin	24 vials
BiotinQuant™ Assay Buffer	25ml
Biotin Standard	1ml

STORAGE CONDITION

The kit is shipped at Ambient Temp. Upon arrival, store the kit components at -20°C. Once the biotin reagent has been opened, store at -20°C with a desiccant as reagent is moisture sensitive. Allow to warm to room temperature before opening.

SPECIFICATIONS

Molecular weight: 556.59 Spacer Arm (Å): 22.4 Membrane Permeable: No

Water Soluble: Yes Reaction pH: 7-9

PRECAUTIONS

- Dissolve the HOOK[™] Sulfo-NHS-LC-Biotin immediately prior to use as the NHS-ester will hydrolyze and become inactive. Do not prepare stock solutions.
- Avoid using primary amine containing buffers, such as Tris and Glycine, as these will compete with the reaction.

ADDITIONAL ITEM(S) REQUIRED

15ml collection tubes

PREPARATION BEFORE USE

- Dilute and prepare 1X Optimizer Buffer™ (1ml 5X Optimizer Buffer™ per 4ml deionized water).
- 2. Warm the Biotin-Agent vial(s) to room temperature before opening to prevent the condensation and deterioration of the biotin agent.
- 3. Add 50µl ultra pure water to a vial of OneQuant™ HABA/Avidin. Incubate at room temperature for 5 minutes. Vortex to solubilize the HABA/Avidin.

A. PROTEIN SAMPLE PREPARATION

- 1a Dissolve 1-10mg protein in 0.5-2ml 1X Optimizer Buffer™ I to a maximum concentration of 10mg/ml.
- 1b If your protein is in an amine—free buffer at a pH of 7.2-8.0 then proceed to the next section.
- 1c For protein in Tris or other amine containing buffers a buffer exchange must be performed. The buffer exchange can be done by dialysis against Optimizer Buffer™ I, we recommend using our Tube-O-DIALYZER™ micro dialysis devices that ensure no loss of precious protein (See Appendix 1). Or one of the supplied Spin-OUT™ columns can be used for buffer exchange as described in Section E. Please note this kit is designed for 10 reactions and the Spin-OUT™ columns are for purification of the biotin labeled protein, using a column for buffer exchange will reduce the number of reactions that can be performed. Additional columns can be ordered at www.GBiosciences.com.

B. CALCULATION OF OUANTITY OF BIOTIN AGENT NEEDED FOR CONJUGATION

To achieve approximately 4-6 biotin groups per antibody molecule, we recommend using a 20 molar excess of biotin to antibody. The extent of biotin labeling for other proteins is dependent on the distribution of amine groups and size of the protein, therefore the molar ratio can be adjusted to suit your needs.

1. Millimoles of HOOK™ Sulfo-NHS-LC-Biotin to be added for a 20 mole excess:

Protein Sample
Volume (ml)

Protein Sample
X Concentration (mg/ml)
Protein Mol. Wt (Da)

X 20 = mmol HOOK™ Sulfo-NHS-LC-Biotin

2. µl 10mM HOOK™ Sulfo-NHS-LC-Biotin to add:

mmol HOOK[™] Sulfo-NHS-LC-
Biotin
$$X = 556.59 \times \frac{500}{2.8} = \frac{\mu l \text{ HOOK}^{\text{™}} \text{ Sulfo-NHS-LC-Biotin}}{\text{solution}}$$

556.59= HOOK™ Sulfo-NHS-LC-Biotin molecular weight

500 = μ l of water 2.8mg of HOOK™ Sulfo-NHS-LC-Biotin dissolved in for a 10mM solution Example: For 0.5ml of a 5mg/ml IgG solution (150,000 Mol. Wt) solution.

0.5ml
$$X = \frac{5mg/ml}{150,000Da} X$$
 20 = 0.000333mmol HOOK[™] Sulfo-NHS-LC-Biotin

0.000333mmol HOOKTM Sulfo-NHS-
LC-Biotin
$$X$$
 556.59 X $\frac{500}{2.8}$ = $\frac{33.1\mu l \text{ HOOK}^{TM} \text{ Sulfo-NHS-LC-}}{\text{Biotin solution}}$

C. PREPARATION OF BIOTIN AGENTS

- 1. Warm the biotin-agent vials to room temperature before opening.
- 2. Immediately before using, add 500µl deionized water to every 2.8mg HOOK™ Sulfo-NHS-LC-Biotin for a 10mM working solution.

NOTE: Make fresh each time and do not prepare stock solutions.

D. BIOTIN CONJUGATION REACTION

- 1. Add the calculated volume (Section B) of freshly prepared 10mM HOOK™ Sulfo-NHS-LC-Biotin to the protein solution from Section A.
- Incubate the reaction at room temperature for 30-60 minutes or on ice for 2 hours.
 Longer incubations can be performed, however this may result in increased chance of protein degradation.

E. REMOVAL OF UNCONJUGATED BIOTIN AGENTS

- Prepare the Spin-OUT[™] column by removing the top and then bottom caps. Place into a 15ml collection tube.
- 2. Centrifuge the column at 1,000g for 2 minutes to remove the storage buffer. Discard storage buffer and return column to 15ml collection tube.
- 3. Equilibrate the column with 2ml 1X Optimizer Buffer™ I, by adding slowly to the resin bed. Centrifuge at 1,000g for 2 minutes. Discard flow through and repeat this step a further 2 times.
- 4. Place the column in to a clean 15ml collection tube and apply the sample directly to the center of the resin bead. Allow the sample to migrate into the resin bed.
- 5. Centrifuge the column at 1,000g for 2 minutes. The flow through is the purified labeled protein sample.
- 6. Store biotinylated protein at 4°C in 0.1% sodium azide until ready for use. Store at -20°C for long term storage.

F. ESTIMATION OF BIOTIN INCORPORATION EFFICIENCY

The method of biotin incorporation estimation is based on the binding of avidin with HABA dye (2-(4-Hydroxyphenylazo)benzoic acid/ 2-(4'-Hydroxybenzeneazo)benzoic acid/ 4'-Hydroxyazobenzene-2-carboxylic acid), which produces a color that can be read at 500nm. The HABA-avidin complex can be displaced with free biotin or biotin conjugated with other molecules (proteins). Measuring the change in optical density of HABA-avidin complex with biotinylated proteins allows for accurate estimation of the molar ration of biotin conjugated to the protein/ antibody.

Important Information

- Ensure that all free/ unconjugated biotin is removed from the labeled protein or
 other molecule before performing an estimation. We recommend desalting with
 our SpinOUT™ desalting spin columns or dialysis with our micro dialysis devices,
 Tube-O-DIALYZER™.
- During desalting or dialysis, we recommend exchanging the reaction buffer to BiotinQuant™ Assay Buffer to ensure accurate estimation. PBS or TBS may also be used, but avoid buffers containing potassium that may result in unwanted precipitation.
- A small variation in color between the OneQuant™ HABA/Avidin does not affect the performance of the reagents.

• The Biotin Standard is supplied as an optional positive control for the assay. Use 100µl in lieu of the biotinylated sample. See calculation for determining amount of biotin in the standard.

Protocol 1: Cuvette Protocol

- 1. Allow the reagents to warm to room temperature.
- 2. Pipette 850µl BiotinQuant™ Assay Buffer into a 1ml cuvette and zero the spectrophotometer at a 500nm wavelength.
- Briefly centrifuge a OneQuant[™] HABA/Avidin vial and then transfer entire contents to the cuvette and mix by gentle inversion.
- 4. Measure the absorbance of the HABA/Avidin complex at 500nm. This is your A_{500} HABA/Avidin reading.
- Add 100µl biotinylated sample to the HABA/Avidin cuvette and mix well by inversion.
 - NOTE: If using optional Biotin Standard, replace the 100µl biotinylated sample with 100µl Biotin Standard.
- 6. Measure the absorbance of the solution at 500nm. Record the absorbance once it has stabilized for 10-15 seconds. This is your A_{500} HABA/Avidin/Biotin Sample reading.
 - NOTE: If the absorbance is <0.3, dilute the biotin sample and repeat the assay.
- 7. Go to the calculation section to determine the moles of biotin per mole of protein.

Protocol 2: Microplate Protocol

- 1. Allow the reagents to warm to room temperature.
- Pipette 170µl BiotinQuant™ Assay Buffer into each microplate well. Blank the plate reader with a well containing only BiotinQuant™ Assay Buffer.
- Briefly centrifuge a OneQuant™ HABA/Avidin vial and then add 10µl OneQuant™ HABA/Avidin to the cuvette and mix on an orbital shaker or equivalent.
- 4. Measure the absorbance of the HABA/Avidin complex at 500nm. This is your A_{500} HABA/Avidin reading.
- 5. Add 20µl biotinylated sample to the HABA/Avidin well and mix well as before. NOTE: If using optional Biotin Standard, replace the 20µl biotinylated sample with 20µl Biotin Standard.
- 6. Measure the absorbance of the solution at 500nm. Record the absorbance once it has stabilized for 10-15 seconds. This is your A_{500} HABA/Avidin/Biotin Sample reading.
 - NOTE: If the absorbance is <0.3, dilute the biotin sample and repeat the assay.
- 7. Go to the calculation section to determine the moles of biotin per mole of protein.

Calculations

Based on Beer Lambert (Beer's) Law: $A_{\lambda} = \varepsilon_{\lambda}bC$, where

- A is the absorbance at a particular wavelength (λ). HOOK™ BiotinQuant™ assay is performed at 500nm.
- ϵ is the extinction coefficient at the wavelength (λ). For HABA/Avidin samples at 500nm, pH7.0 this is 34,000M⁻¹cm⁻¹.
- **b** is the path length in centimeters. Cuvettes (10x10mm) have a pathlength of 1cm. The pathlength for microplates, using the indicated volumes, is normally 0.5cm.
- C is the molarity concentration of the sample (= mol/L = mmol/ml)

For calculating the number of moles of biotin per mole of protein or sample the following values are required:

- Concentration of protein/sample used (mg/ml)
- Molecular weight of protein, expressed as grams per mole (e.g. IgG = 150,000)
- A₅₀₀ HABA/Avidin reading
- A₅₀₀ HABA/Avidin/Biotin Sample
- Dilution factor (DF), if sample was diluted before adding to HABA/avidin solution.
- 1. Calculate mmol biotinylated protein/ml:

2. Calculate change in absorbance at 500nm:

Calculation #2	(0.9 x A ₅₀₀ HABA/Avidin) –(A ₅₀₀	- A A
(Cuvette):	HABA/Avidin/Biotin Sample)	=ΔA ₅₀₀

Calculation #2
$$(A_{500} \text{ HABA/Avidin}) - (A_{500} \text{ HABA/Avidin/Biotin}$$

= ΔA_{500}

(Microplate): Sample)

NOTE: 0.9 is the correction factor for the dilution of the HABA/Avidin with the sample in the cuvettes. This is not necessary for microplates as the dilution is offset by the increase in volume and therefore the light path (b).

3. Calculate concentration of biotin in reaction (mmol/ml):

Calculation #3:
$$\frac{\Delta A_{500}}{34,000 \times b} = \frac{\text{Calculation #2}}{34,000 \times b} = \frac{\text{mmol biotin}}{\text{ml reaction mixture}}$$

NOTE: b = lightpath, which is 1cm for cuvettes and 0.5cm for microplates.

4. Calculate mmol of biotin per mmol of protein:

	mmol biotin in		mmol biotin in		Calculation #3 x
Calculation #	t original sample		reaction x 10 x DF		10 x DF
4:	mmol protein in	-	Calculation #1	•	Calculation #1

NOTE: DF is the dilution factor. 10 is for the 10 fold dilution of the biotinylated protein sample in the reaction mixture.

5. Calculate concentration of biotin in Biotin Standard (mM):

Calculation #5:
$$\frac{\Delta A_{500} \times 10 \times 1000}{34,000 \times b} = [Biotin Standard] (mM)$$

NOTE: b = lightpath, which is 1cm for cuvettes and 0.5cm for microplates. 10 is for the 10 fold dilution of the Biotin Standard in the reaction mixture.

TROUBLESHOOTING

Issue	Suggested Reason	Possible Solution		
	Low or zero biotinylation of protein.	Lack of functional groups for biotinylation, use a different coupling chemistry.		
ΔA ₅₀₀ is ≤0	Incomplete reagent mixing	Ensure all the OneQuant™ HABA/Avidin is fully dissolved before using		
	Particulates in protein solution interfering with absorbance	Filter protein solution before assaying		
	Potassium ions present in sample	Ensure samples are in BiotinQuant™ Assay Buffer		
Biotin levels are unexpectedly high	Free, Unconjugated biotin not removed	Desalt or dialyze biotinylated sample before use to remove free biotin.		

APPENDIX 1: SAMPLE EQUILIBRATION WITH TUBE-O-DIALYZER™ (NOT SUPPLIED)

If protein solution is in an incompatible buffer, dialyze and equilibrate into 1X Optimizer Buffer™ as follows:

- Pipette your sample directly into the Tube-O-DIALYZER™ tube. For Tube-O-DIALYZER™ Micro use 20-250µl and for Tube-O-DIALYZER™ Medi use 0.2-2.5ml.
 NOTE: Tube-O-Dialyzer™ is available in 1, 4, 8, 15 and 50kDa MWCO. Visit our website for further information.
- Pipette 3-5ml appropriate 1X Optimizer Buffer™ into a Micro Dialysis Cup or small beaker. If a small magnetic stir bar is available add to the Micro Dialysis Cup, if not add 3-5 glass balls.
- Screw the dialysis cap on to the Tube-O-DIALYZER™ tube. Invert the Tube-O-DIALYZER™, ensuring the entire sample rests upon the membrane.
 NOTE: If sample is too viscous, centrifuge the Tube-O-DIALYZER™ in an inverted position (i.e. the dialysis membrane facing downward). Centrifuge for 5 seconds at 500-1,000q.
- 4. Keeping the Tube-O-DIALYZER™ in an inverted position, slide the supplied float onto the Tube-O-DIALYZER™ tube. Place the Tube-O-DIALYZER™ in the Micro Dialysis Cup with the Optimizer Buffer™.
- 5. Ensure that the dialysis membrane contacts the dialysis buffer. If there are large air bubbles trapped underneath the dialysis membrane surface, tilt the tube or squirt buffer to remove the air bubbles. Gently, stir the dialysis buffer with a magnetic stir or place on an orbital shaker. For efficient and complete dialysis we recommend inverting or gently tapping the Tube-O-DIALYZER™ 1-2 times during dialysis to mix the sample. If necessary repeat the centrifugation in step 3.
- 6. Dialyze at room temperature, or 4°C if required, for 1-2 hours.
- 7. Repeat the dialysis with 1-2 changes of buffer.
- 8. After dialysis, remove the Tube-O-DIALYZER™ from the float and immediately spin the Tube-O-DIALYZER™ (in up-right position) for 5-6 seconds at 500-1,000xg.

APPENDIX 2: INSTRUCTIONS FOR CELL SURFACE PROTEIN BIOTINYLATION

NOTE: For cell surface protein labeling, we recommend our HOOK™ Cell Surface Protein Isolation kit that uses G-Biosciences HOOK™ biotin labeling and purification technology in conjunction with our Mammalian Cell PE LB™ lysis buffer to conveniently label cell surface proteins and isolate them for further analysis, including Western blotting.

Cell Sample Preparation:

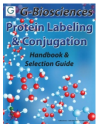
- Wash cells three times with ice-cold PBS buffer to remove any contaminating proteins.
- Suspend the cells at a concentration of 25 x 10⁶ cells/ml in PBS.
 NOTE: Other cell concentrations can be used based on cell size, type, etc. The concentration of biotinylation reagent can be scaled up or down accordingly.

Biotin Agent & Cell Surface Reaction

- Prepare HOOK™ Sulfo-NHS-LC-Biotin reagent by adding as described above 0.5mg
 Biotin Agent per ml of reaction volume (solid form of biotin agent can be directly
 added if it is water soluble).
- 2. Gently mix and incubate at room temperature for 30 minutes.
- 3. Wash cells three times with ice-cold PBS to remove any remaining biotinylation reagent. The cells surface proteins are now biotinylated.

RELATED PRODUCTS

Download our Protein Labeling & Conjugation Handbook



http://info.gbiosciences.com/complete-protein-labeling-conjugation-handbook/
For other related products, visit our website at www.GBiosciences.com or contact us.

Last saved: 2/4/2019 CMH



www.GBiosciences.com