# **INSTRUCTIONS**



0418.8

# DSS and BS<sup>3</sup> Crosslinkers

# <u>21555 21580 21585 21586 21655 21658</u>

Description	
DSS (disuccinimidyl suberate), 1g	
<b>DSS</b> , 50mg	
DSS, No-Weigh <sup>TM</sup> Format, 8 × 2mg microtubes	
Molecular Weight: 368.35	
Spacer Arm: 11.4Å	0
Formula: $C_{16}H_{20}N_2O_8$	
<b>BS</b> <sup>3</sup> (bis[sulfosuccinimidyl] suberate), 1g	
<b>BS</b> <sup>3</sup> , 50mg	Na <sup>+</sup> O <sup>-</sup>
<b>BS<sup>3</sup></b> , No-Weigh Format, 8 × 2mg microtubes	
Molecular Weight: 572.43	0 0 0 0 Na <sup>+</sup>
Spacer Arm: 11.4Å	0
Formula: $C_{16}H_{18}N_2Na_2O_{14}S_2$	
	DescriptionDSS (disuccinimidyl suberate), 1gDSS, 50mgDSS, No-WeighTM Format, $8 \times 2mg$ microtubesMolecular Weight: 368.35Spacer Arm: 11.4ÅFormula: $C_{16}H_{20}N_2O_8$ BS³ (bis[sulfosuccinimidyl] suberate), 1gBS³, 50mgBS³, No-Weigh Format, $8 \times 2mg$ microtubesMolecular Weight: 572.43Spacer Arm: 11.4ÅFormula: $C_{16}H_{18}N_2Na_2O_{14}S_2$

Storage: Upon receipt store product desiccated at 4°C. Products are shipped at ambient temperature.

### Introduction

Thermo Scientific DSS is a water-insoluble, homobifunctional *N*-hydroxysuccinimide ester (NHS ester), and BS<sup>3</sup> is its watersoluble analog. NHS esters react efficiently with primary amino groups  $(-NH_2)$  in pH 7 to 9 buffers to form stable amide bonds. The reaction results in the release of *N*-hydroxysuccinimide. Proteins, including antibodies, generally have several primary amines in the side chain of lysine (K) residues and the N-terminus of each polypeptide that are available as targets for NHS-ester reagents.

The water-soluble and insoluble forms of NHS-esters have essentially identical reactivity toward primary amines. BS<sup>3</sup> is supplied as a sodium salt and is water-soluble up to 10mM. DSS is hydrophobic and dissolved in an organic solvent such as DMSO or DMF then added to the aqueous reaction mixture. DSS does not possess a charged group and is lipophilic and membrane-permeable, which makes it useful for intracellular and intramembrane conjugations. Water-soluble BS<sup>3</sup> possess a charged group and is useful for cell-surface protein crosslinking.

# **Important Product Information**

- DSS and BS<sup>3</sup> are moisture-sensitive. To avoid moisture condensation onto the product, vial must be equilibrated to room temperature before opening.
- Prepare these crosslinkers immediately before use. The NHS-ester moiety readily hydrolyzes and becomes non-reactive; therefore, do not prepare stock solutions for storage. Discard any unused reconstituted crosslinker.
- No-Weigh Microtube Handling: Each of the eight microtubes contains 2mg of crosslinker. Puncture foil with a pipette tip and add DMSO or DMF to the DSS or aqueous buffer to the BS<sup>3</sup>. Store the microtube strip in the foil pouch provided. Used microtubes may be cut from the unused microtubes and discarded.
- Crosslinking proteins with biological activity (i.e., enzymes, antibodies etc.) can result in activity loss upon conjugation possibly caused by conformational changes of the molecule. Activity loss also may occur when the crosslinker modifies lysine groups involved in binding substrate or antigen. Adjusting the molar ratios of reagent to the target may overcome activity loss. Alternatively, use a crosslinker that targets a different functional group.



• Hydrolysis of the NHS ester is a competing reaction and increases with increasing pH. Hydrolysis occurs more readily in dilute protein or peptide solutions. In concentrated protein solutions, the acylation reaction is favored.

## **Procedure for Crosslinking Proteins**

The following protocol is an example application for this product. Specific applications will require optimization.

#### A. Materials Required

- Dry dimethyl sulfoxide (DMSO) or dimethyl formamide (DMF) for use with DSS
- Conjugation Buffer: Use a non-amine-containing buffer at pH 7 to 9, such as 100mM sodium phosphate, 0.15M NaCl (Product No. 28372); 20mM HEPES; 100mM carbonate/bicarbonate; or 50mM borate
- Quenching Buffer: 1M Tris•HCl, pH 7.5 (1M glycine or lysine also may be used) non-reacted reagent can be removed by dialysis or gel filtration.

#### **B.** Procedure

- 1. Prepare protein in Conjugation Buffer.
- 2. Prepare crosslinker immediately before use. Dissolve BS<sup>3</sup> first in water or 20mM sodium phosphate buffer, as more concentrated buffer salt may interfere with initial solubility of the reagent; once the BS<sup>3</sup> is dissolved, the solution can be diluted or added to more concentrated buffer solutions without adversely affecting its solubility. Prepare DSS by dissolving in DMSO or DMF. Examples for preparations are as follows:
  - Microtubes: Each No-Weigh Microtube contains 2mg of crosslinker.
  - **Powder:** Weigh 2mg of crosslinker into a microcentrifuge tube.

<u>Solve</u> add	nt volume to to 2mg DSS	Buffer volume to add to 2mg BS <sup>3</sup>	Crosslinker Concentration
	432μL*	277μL*	12.5mM
	216µL	140µL	25mM
	108μL	70µL	50mM
	54µL	35µL	100mM

\*The maximum volume that can be added to the No-Weigh Microtube is  $220\mu$ L.

- 3. Add crosslinker to the protein sample. If the protein concentration is greater than 5mg/mL, use a 10-fold molar excess of the crosslinker. For samples < 5mg/mL, use a 20- to 50-fold molar excess of the crosslinker. Use a final concentration of crosslinker at 0.25-5mM.
- 4. Incubate the reaction mixture at room temperature for 30 minutes or on ice for 2 hours.
- 5. Quench the reaction using by adding Quenching Buffer to a final concentration of 20-50mM Tris. Alternatively, remove the non-reacted reagent by dialysis or desalting.
- 6. Incubate the quenching reaction at room temperature for 15 minutes.

# Procedure for Intra- and Extracellular Crosslinking

Crosslinking may be performed on cells in suspension or on adherent cells in culture plates. In the latter situation, diffusion of the crosslinking reagent to all surfaces of the cells will be limited and will occur predominately on the exposed surface. Culture media must be washed from the cells otherwise amine-containing components will quench the reaction. Using a more concentrated cell suspension is most effective as less reagent will be required in the reaction. Generally, a final concentration of 1-5mM reagent is effective. NHS reactions occur more rapidly with increasing pH; therefore, pH 8.0 is used in the following example so the reaction can be completed quickly.

**Note:** Use membrane-insoluble BS<sup>3</sup> for crosslinking molecules on the cell surface. Use DSS when crosslinking within the cell is required.



#### A. Materials Required

- Crosslinker Solution: Immediately before use, dissolve the DSS in dry DMSO at 10-25mM. Dissolve BS<sup>3</sup> in water or buffer. BS<sup>3</sup> may be added directly to the cells to decrease the extent of hydrolysis.
- Phosphate-buffered saline (PBS): 20mM sodium phosphate, 0.15M NaCl; pH 8. HEPES, bicarbonate/carbonate or borate buffers between pH 7 and 9 may be used as alternative buffers.
- Quench Solution: 1M Tris, pH 7.5 (Tris or glycine will quench the reaction.)

#### B. Procedure

- 1. Suspend cells at  $\sim 25 \times 10^6$  cells/mL in PBS (pH 8.0).
- Wash cells three times with ice-cold PBS (pH 8.0) to remove amine-containing culture media and proteins from the cells. Note: For cell-surface interaction studies, add ligands to the cells and incubate for 1 hour at 4°C.
- 3. Add the DSS or  $BS^3$  solution to a final concentration of 1-5mM.
- Incubate the reaction mixture for 30 minutes at room temperature.
  Note: Performing this incubation at 4°C may reduce active internalization of BS<sup>3</sup>.
- 5. Add the Quench Solution to a final concentration of 10-20mM Tris.
- 6. Incubate the quenching reaction for 15 minutes at room temperature.

# **Related Thermo Scientific Products**

20036	Bioconjugate Techniques, 1202 pages, softcover
66382	Slide-A-Lyzer <sup>®</sup> Dialysis Cassette Kit, 10K MWCO, 3mL, 10 cassette kit
66807	Slide-A-Lyzer Dialysis Cassette Kit, 10K MWCO, 12mL, 8 cassette kit
22585	DSP (dithiobis[succinimidylpropionate]), 1g, cleavable NHS-ester crosslinker
21578	DTSSP (3,3'-dithiobis[sulfosuccinimidylpropionate]), 50 mg, cleavable Sulfo-NHS-ester crosslinker
28372	BupH <sup>TM</sup> Phosphate Buffered Saline Packs, 40 pack

#### **General References**

Cox, G.W., et al. (1990). Characterization of IL-2 receptor expression and function on murine macrophages. J Immunol 145:1719-26.

Knoller, S., *et al.* (1991). The membrane-associated component of the amphiphile-activated, cytosol-dependent superoxide-forming NADPH oxidase of macrophages is identical to cytochrome b559. *J Biol Chem* **266**:2795-2804.

Partis, M.D., et al. (1983). Cross-linking of protein by ω-maleimido alkanoyl N-hydroxysuccinimido esters. J Prot Chem 2(3):263-77.

This product ("Product") is warranted to operate or perform substantially in conformance with published Product specifications in effect at the time of sale, as set forth in the Product documentation, specifications and/or accompanying package inserts ("Documentation") and to be free from defects in material and workmanship. Unless otherwise expressly authorized in writing, Products are supplied for research use only. No claim of suitability for use in applications regulated by FDA is made. The warranty provided herein is valid only when used by properly trained individuals. Unless otherwise stated in the Documentation, this warranty is limited to one year from date of shipment when the Product is subjected to normal, proper and intended usage. This warranty does not extend to anyone other than the original purchaser of the Product ("Buyer").

No other warranties, express or implied, are granted, including without limitation, implied warranties of merchantability, fitness for any particular purpose, or non infringement. Buyer's exclusive remedy for non-conforming Products during the warranty period is limited to replacement of or refund for the non-conforming Product(s).

There is no obligation to replace Products as the result of (i) accident, disaster or event of force majeure, (ii) misuse, fault or negligence of or by Buyer, (iii) use of the Products in a manner for which they were not designed, or (iv) improper storage and handling of the Products.

Current product instructions are available at <u>www.thermoscientific.com/pierce</u>. For a faxed copy, call 800-874-3723 or contact your local distributor. © 2011 Thermo Fisher Scientific Inc. All rights reserved. Unless otherwise indicated, all trademarks are property of Thermo Fisher Scientific Inc. and its subsidiaries. Printed in the USA.