

Sample Preparation Kit (96 reactions)

PROTOCOL – CHEMICAL LABELLING



INTRODUCTION

Sample preparation is one of the essential steps of bottom-up proteomics. The PreOmics NHS-compatible sample preparation kit is designed to assist you achieving best results with few sample preparation steps and little hands-on time for multiplexing applications. For sample specific protocols and optimization contact us or visit our website at www.preomics.com.

KIT CONTENTS

The iST-NHS kit provides a streamlined solution for reliable sample preparation compatible with chemical labelling such as iTRAQ or TMT. It includes all chemicals to denature, reduce and alkylate proteins, as well as the enzymes to perform a tryptic digest and a final clean-up.

COMPONENT	CAP COLOR	QUANTITY	BUFFER PROPERTIES				DESCRIPTION	STORAGE
			Organic	Acidic	Basic	Volatile		
DIGEST	red	24x					Enzyme <i>Trypsin</i> -mix to digest proteins.	-20 °C
RESUSPEND	yellow	4x 2 mL				✓	Protease reconstitution buffer for enzymes.	RT
LYSE-NHS	orange	12x 1 mL				✓	Denature, reduce and alkylate proteins.	RT
STOP	black	12x 1 mL	✓	✓		✓	Stop the enzymatic activity.	RT
WASH 1	blue	12x 2 mL	✓	✓		✓	Clean up peptides from hydrophobic contaminants.	RT
WASH 2	green	12x 2 mL				✓	Clean up peptides from hydrophilic contaminants.	RT
ELUTE	violet	12x 2 mL	✓			✓	Elute the peptides from the cartridge.	RT
LC-LOAD	white	12x 1 mL				✓	Load peptides on reversed-phase LC-MS column.	RT
CARTRIDGES		96x					Cartridges for 1 to 100 µg protein starting material.	RT
WASTE PLATE		1x					Deep well plate for collecting waste after washes.	RT
MTP PLATE		1x					LoBind plate for collecting peptides after elution.	RT
ADAPTER PLATE		1x					Enables placing cartridges in the 96well MTP plate.	RT
ADAPTER		8x					Enables placing a cartridge into a tube.	RT
CAPS		96x					Caps to optionally close the cartridge's bottom.	RT

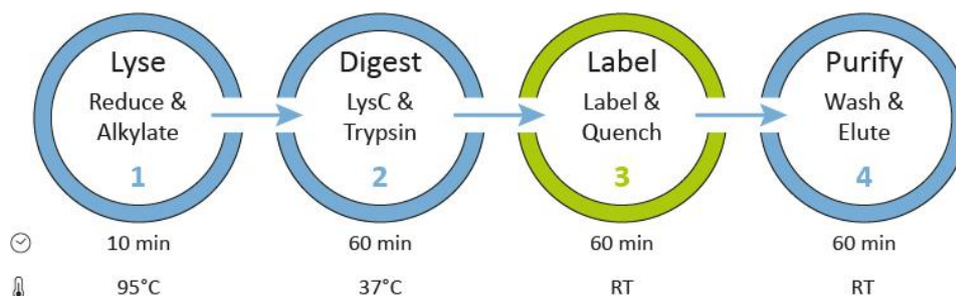
PRE-REQUISITES

Common lab equipment is required for the sample preparation.

EQUIPMENT	DESCRIPTION
PIPETTE	Careful sample handling and pipetting reduces contaminations and improves quantification.
SAMPLE	Pelleted cells or precipitated protein. For other materials ask us for adapted protocols.
96x-WELL	96 deep well and PCR plates with skirt to balance WASTE & MTP plates in centrifuge.
HEATING BLOCK*	2x MTP plate heaters are recommended to help protein denaturation and during digestion.
CENTRIFUGE	Wing-bucket MTP plate centrifuges are necessary for loading, washing and elution.
SONICATOR	If you have DNA in your sample, shear it by sonication (e.g. Diagenode Bioruptor®).
SPEED-VAC	Vacuum manifolds evaporate volatile buffers from the eluate before LC-MS.
ULTRASONIC BATH	Optional: can be used to resuspend peptides.
LABELING REAGENT	Labelling reagent (e.g 400 µg labeling reagent in 41 µL dry acetonitrile for 100 µg peptides).
LABELING BUFFERS	Dry acetonitrile & quenching buffer (5% hydroxylamine) as recommended by the manufacturer.

* a heating shaker adapter for MTP plate shakers can be purchased from PreOmics

PROCEDURE



Please refer to www.preomics.com for Limited Use Label License and Product Warranty.

Material: Pelleted cells and precipitated protein
 Quantity: 1 - 100 µg protein starting material

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PROTOCOL

1. LYSE

- 1.1. Add 50 µL **LYSE-NHS** (orange dot) to 1-100 µg of protein sample, place it in a HEATING BLOCK (95 °C; 1,000 rpm; 10 min).
- 1.2. Optional: Spin down droplets (RT; max. 300 rcf; 10 sec). ***NOTE 1***
- 1.3. If you expect DNA, shear sample in a SONICATOR (10 cycles; 30 sec ON/OFF).
- 1.4. Transfer sample to **CARTRIDGE** and cool down (RT). Be careful not to damage the bottom layer of **CARTRIDGE**.
- 1.5. Use **ADAPTER PLATE** to place **CARTRIDGES** on top of the **WASTE PLATE** tube. Label all plate and wells.

2. DIGEST

- 2.1. Add 210 µL **RESUSPEND** (yellow dot) to **DIGEST** (red dot) (1 tube for 4 reactions), shake (RT; 500 rpm; 10 min), pipette up/down.
- 2.2. Add 50 µL **DIGEST** (red dot) to **CARTRIDGE** and place it in a pre-heated HEATING BLOCK (37 °C; 500 rpm; 1-3 hours).
- 2.3. Resuspend LABELING REAGENT in dry acetonitrile (see page 1; 4:1 ratio of label:peptides). ***NOTE 2***
- 2.4. Add resuspended LABELING REAGENT to **CARTRIDGE**, pipette up/down, incubate shaking (RT; 500 rpm; 1 hour).
- 2.5. Add 10 µL QUENCHING BUFFER (5% Hydroxylamine) to **CARTRIDGE** and mix.
- 2.6. Add 100 µL **STOP** (black dot) to **CARTRIDGE** (precipitation may happen), shake (RT; 500 rpm; 1 min), pipette up/down. ***SP***

3. PURIFY

- 3.1. Spin **CARTRIDGE** in a CENTRIFUGE (3,800 rcf; 1-3 min). If needed, adjust values to ensure complete flow-through.
- 3.2. Add 200 µL **WASH 1** (blue dot) to **CARTRIDGE**, repeat step 3.1., discard flow-through.
- 3.3. Add 200 µL **WASH 2** (green dot) to **CARTRIDGE**, repeat step 3.1., discard flow-through. ***SP***
- 3.4. Use **ADAPTER PLATE** to place **CARTRIDGES** on top of the **MTP PLATE**. Label plate and wells.
- 3.5. Add 100 µL **ELUTE** (purple dot) to **CARTRIDGE**, repeat step 3.1., keep flow-through in **MTP PLATE**.
- 3.6. Repeat step 3.5., keep flow-through in the same **MTP PLATE**.
- 3.7. Remove **CARTRIDGES** and place **MTP PLATE** in a SPEED-VAC (45 °C; until completely dry).
- 3.8. Add **LC-LOAD** (white dot) to **MTP PLATE**. Aim for 1 g/L concentration (e.g. 100 µL to 100 µg protein starting material).
- 3.9. Sonicate **MTP PLATE** in an ULTRASONIC BATH (5 min) or shake (RT; 500 rpm; 5 min). ***SP***

NOTE 1 Volumes of buffers can be adjusted according to protein starting amounts. Lysis temperature should be between 60-95 °C. Most classical protein quantification assays are compatible with our lysis buffer. We recommend the BCA assay or tryptophan quantification method. Some assays require dilution with water to achieve best results (Dilutions: BCA: none; Bradford: 1:4; Coomassie: 1:20; Lowry: 1:4; Tryptophan: none). Visit our FAQ website for more information.

NOTE 2 We recommend to use only fresh labeling reagents at a label to peptide ratio of 4:1, as well as a min. 30% (v/v) acetonitrile concentration during the labeling reaction to achieve best labeling and identification results. In case of reduced labeling efficiency, use LABELING REAGENT concentrations according to the manufacturer's instructions and see the manufacturer's troubleshooting information.

***SP* - Storage Point:** At this point, close the peptide containing tube or **CARTRIDGE** (use a **CAP** for bottom). Peptides can be frozen at -20 °C. Storage of peptides should not exceed 2 weeks at -20 °C. For extended storage, finish the protocol and store at -80 °C.

DATA ANALYSIS

Consider the following as fixed modifications in your database search:

MODIFICATION	DESCRIPTION	COMPOSITION	SPECIFICITY	MASS
ALKYLATION	Specific cysteine modification	C ₆ H ₁₁ NO	[C]	+113.084 Da