### **Product Information**

# **Cytoplasmic and Nuclear Protein Extraction Kit**

## Catalog Number: NC-001

#### **Description:**

This kit is designed for extracting intact nuclear proteins and native, non-denatured cytoplasmic proteins from various cell types or tissues, prepared for EMSA, ELISA, 1D and 2D electrophoresis, Western blotting, TF-TF interaction arrays and other protein/DNA assays.

#### Kit contains:

Components	Quantity (50 extractions)	Storage
Cytoplasmic Lysis Buffer (C207020, Blue sticker)	25.0 mL	2-8°C
Cytoplasmic Washing Buffer (C207030 Purple sticker)	15.0 mL	2-8°C
Detergents (D207050 Yellow cap)	1.5 mL	2-8°C
Nuclear Lysis Buffer (N207040, Green sticker)	2.5 mL	2-8°C
DTT, 1M (Dissolved in 0.1 ml ddH2O)	1 vial	-20°C
Protease/Phosphatase Inhibitors (I208052) supplied in DMSO, contains optimized AEBSF, Aprotinin, E64, Leupeptin, Per	1  vial pstatin A, Sodium fluoride, Sodium orthovanadate and S	-20°C Sodium pyrophosphate.

# **Protocol:** (Keep all buffers and cell/tissue samples on ice)

\*Prepare working reagents prior to proceeding.

For 10 Extractions: (10e7 cells or 50 mg tissues/Extraction)		
Cytoplasmic Lysis Buffer (5ml)	add 5.0ul (1M DTT ) and 100ul Protease/phosphatase Inhibitors $_{\scriptscriptstyle (I208052)}$	
Cytoplasmic Washing Buffer (3.0ml)	add 2.0ul (1M DTT ) and 30ul Protease/phosphatase Inhibitors (1208052)	
Nuclear Lysis Buffer (0.5ml)	add 0.5ul (1M DTT ) and 20ul Protease/phosphatase Inhibitors (I208052)	

## 1. Preparation of samples from culturing/frozen cells:

Harvest cells (1x 10e7 cells) as usual and wash cells once with 1.0ml 1x ice-cold PBS/DPBS, centrifuge at 1,600 rpm for 8 minutes, aspirate liquids. Add <u>500ul cytoplasmic lysis buffer</u> to resuspend cell pellet. Gently pipette up and down several times and incubate on ice for 10 minutes.

## Preparation of samples from tissues:

 Weigh 10-50mg frozen/ fresh tissues and chop tissues into small pieces using a clean razor blade. Immediately transfer into a 2.0ml microcentrifuge tube contained <u>500ul cytoplasmic</u> <u>lysis buffer</u>. Vortex at mid-speed for 20 seconds and incubate on ice for 10 minutes. Tissues homogenization:

1) Using a clean pre-chilled Teflon pestle homogenizer to homogenize the tissues for 10-20 strokes on ice, simply spin down the cells/tissue suspension and continue to homogenize tissues another 10-20 strokes.

2) (**Alternative-1**): Prepare a syringe with a needle gauged between 23 and 25. Pass cells/tissues through needle about 20 times to disrupt the cell membrane and release the intact nuclei and organelles.

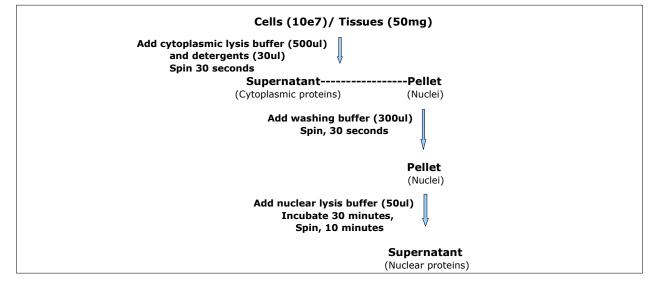
3) (**Alternative-2**): Using a pre-chilled, clean Dounce homogenizer to homogenize the cells/tissues twice at speed 4 (moderate) speed for 20 seconds.

- 2. Add <u>30ul of detergents (yellow cap)</u>, vortex vigorously at highest speed for 10 seconds.
- 3. Centrifuge at 14,000 xg for 30 seconds at 4°C, immediately transfer the supernatant (cytoplasmic protein fractions) into a pre-chilled micocentrifuge tube.
- 4. Add <u>300ul cytoplasmic washing buffer</u> to resuspend the pellet. Centrifuge at 14,000 xg for 30 seconds at 4°C. Aspirate liquids. (The remained cytoplasmic fractions were washed out).
- 5. Resuspend the pellet in <u>50ul nuclear lysis buffer</u> and vortex vigorously for 10 seconds. Incubate suspension for 30 minutes on ice (vortex 10 seconds every 10 minutes).

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- 6. Centrifuge at 14000xg for 10 minutes at 4°C. Transfer the supernatant (nuclear protein fractions) into a clean pre-chilled 1.5ml microcentrifuge tube.
- 7. Determine the protein concentration of cytoplasmic and nuclear with spectrometers, by Bradford or by BCA Assay. Store all the extracts aliquots at -80°C.

#### Flow Chart of Protein Extraction:



#### Additional information:

- The nuclear protein markers: Lamin B (68kDa), LaminA/C (70 KDa), HDAC, Histone H1 (33KDa), Histon H4(43KDa);
- The cytoplasmic protein markers: GAPDH, anti-b-actin;
- The membrane protein markers: EGFR, Na+/K+ ATPase, anti-Sp1;
- The cytoskeleton protein markers: Vimentin.
- The lysosome protein markers: LAMP1/2/3. Capthepsin D.
- The peroxisome protein markers: PMP70.
- The Zmtech protease/phosphatase Inhibitors (I208052) supplied in DMSO, contains optimized AEBSF, Aprotinin, E64, Leupeptin, Pepstatin A, Sodium fluoride, Sodium Orthovanadate and Sodium pyrophosphate.

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