

Minute[™] Plasma Membrane Protein Isolation and Cell Fractionation Kit

Catalog number: SM-005

Description

The MinuteTM kit represents the next generation product for plasma membrane (PM) isolation and cell fractionation, offering unsurpassed convenience and consistency by eliminating uncontrollable variations associated with traditional homogenization and density gradient centrifugation including two phase partitioning.

How it works: Cells/tissues are first sensitized by buffer A before passing through the proprietary filter in a zigzag manner when high-speed centrifugal force is applied, resulting in a cell lysate containing ruptured cell membranes and intact nuclei. As a result, the nuclear contaminations are virtually eliminated. PM is further separated from the cell lysate (a mixture of crude membranes, intact nuclei, cytosol proteins and organelles) by subsequent differential and density centrifugation with a regular tabletop microcentrifuge. 5 distinct cell fractions (total membrane, PM, cytosol, nucleus and organelles) can be obtained at the completion of the protocol. The procedure can be completed in less than 45 minutes.

Applications

The kit is designed to rapidly isolate native membrane proteins from cultured cells or tissues for applications such as SDS-PAGE, immunoblottings, ELISA, IP, membrane protein structure analysis, 2-D gels, enzyme activity assays and other applications. This kit provides the most rapid method currently available for preparation of native membrane proteins.

Kit components (50T):

- 1. 25 ml buffer A
- 2. 10 ml buffer B
- 3. 50 protein extraction filter cartridges
- 4. 50 collection tubes with cap
- 5. 2 plastic rods
- 6. Tissue dissociation beads

Kit components (4T):

- 1. 2.0 ml buffer A
- 2. 1.0 ml buffer B
- 3. 4 protein extraction filter cartridges
- 4. 4 collection tubes with cap
- 5. 1 plastic rods
- 6. Tissue dissociation beads

Storage: Store Buffer A and Buffer B at -20°C upon arrival.

Additional Materials Required

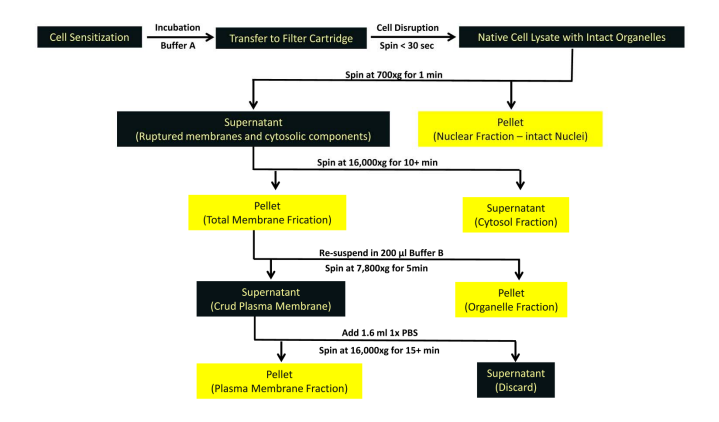
1 X PBS Vortexer Table-Top Microcentrifuge



Important Information:

- 1. Read the entire procedures carefully. Thaw buffer A and buffer B completely, invert the bottles a few times and place them on ice. Chill protein extraction filter cartridge with collection tube on ice prior to use.
- 2. All centrifugation steps should be performed at 4°C in a cold room or in a refrigerated mirocentrifuge.
- 3. To study protein phosphorylation, **phosphatase inhibitors** (such as PhosStop from Roche) should be added to buffer A prior to use. The use of protease inhibitor cocktails is optional.
- 4. It is recommended to use BCA Protein Assay Kit for determination of protein concentration (Pierce, Cat #:23227).

The Workflow





Membrane Protein Isolation Procedures:

A. Isolation of Total Membrane Proteins

- 1. Place the filter cartridges in collection tubs and incubate on ice.
- 2. For cultured cells, collect 1-50 X 10⁶ cells by low speed centrifugation (500-600 X g 5 min). Go to step 3a. For tissue samples, go to step 3b.

Note: For isolation of plasma membrane proteins from cultured cells (see below) it's recommended to use 20-50 X 10⁶ cells.

- 3a. Wash cells once with cold PBS. Remove supernatant completely and resuspend the pellet in buffer A (200 μl for a starting cell number less than 5 million and 500 μl for a starting cell number greater than 5 million). Incubate the cell suspension on ice for 5-10 min. Vortex the tube vigorously for 10-30 seconds. Immediately transfer the cell suspension to the filter cartridge. Go to step 4.
- 3b For tissue samples place a piece of fresh tissue (10-30 mg) or frozen tissue (20-30 mg) in a filter cartridge. Add 200 µl buffer A to the filter and grind the tissue with a plastic rod for one min by pushing the tissue against the surface of the filter repeatedly with twisting force (Note: if you are working with skeletal or cardiac muscles, it is recommended to add 100-120 mg tissue dissociation beads to the filter prior to grinding). Add 300 µl buffer A to the same filter cartridge, mix by pipette up and down a few times and incubate the tube on ice with cap open for 5 min. Go to step 4.

Note: The presence of a small amount of un-homogenized tissue will not affect the quality of the sample. The plastic rod is reusable. For cleaning, wipe it with 75% alcohol or rinse it with distilled water.

4. Cap the filter cartridge and centrifuge at 16,000 Xg for 30 seconds (it is recommended to use a table top centrifuge that can reach maximum speed in less than 10 seconds).

Optional: For cultured cells it is recommended to resuspend the pellet in collection tube from step 4, transfer the cell suspension to the same filter and spin at 16,000Xg for 30 seconds. Re-passing the cells through the filter can increase the yield by 20-30%.

5. Discard the filter and resuspend the pellet by vigorously vortexing for 10 seconds.

Following procedures separate total cellular components into four fractions: nuclei, cytosol, organelles and plasma membrane.

6. Centrifuge at 700 Xg for one min (the pellet contains intact nuclei). Transfer the supernatant to a fresh 1.5 ml microcentrifuge tube and centrifuged at 4°C for 10-30 min at 16,000 X g (longer centrifugation time will increase yield). Remove the supernatant (this is the cytosol fraction) and save the pellet (this is the total membrane protein fraction including organelles and plasma membranes). Store the pellet at -70°C or dissolve it in detergent-containing buffers of your choice. The yield is typically 10-500 µg/sample. You may stop here if isolation of plasma membrane proteins is not needed. Continue to step 7 for plasma membrane



protein isolation. Don't freeze total membrane protein fraction if further isolation of plasma membrane proteins is desired.

B. Isolation of Plasma Membrane Proteins

- 7. Resuspend the total membrane protein fraction from step 6 in 200 µl buffer B by repeatedly pipetting up and down or vortexing. Centrifuge at 7,800 X g for 5 min at 4°C (Note: if final plasma membrane prep is contaminated by organelle membranes, increase centrifugation time up to 20 min can improve the purity). The pellet contains organelle membrane proteins.
- 8. Carefully transfer the supernatant to a fresh 2.0 ml microcentrifuge tube and add 1.6 ml cold PBS. Mix by inverting the tube a few times. Centrifuge at 16,000 X g for 15-30 min (longer centrifugation will improve yield). Discard the supernatant and save the pellet (isolated plasma membrane proteins). Typically, 10-300 µg plasma membrane proteins can be obtained. Pellet of plasma membrane proteins can be dissolved in 20-200 µl detergent containing buffers of your choice depending upon specific downstream applications. Reagents in following table are recommended for solubilization of the pellet. For isoelectric focusing (First dimension of 2D gel) we recommend to use: 7M urea/2M thio-urea/2% Chaps and 20 mM DTT (add DTT to above mix prior to use).

Following protein solubilization reagents are recommended.

| Product Name | Cat. No. | Applications |
|--|----------|---|
| Minute TM Denaturing Protein Solubilization Reagent | WA-009 | SDS-PAGE electrophoresis and Western blotting, trypsin digestion, purification of proteins with biotin labeling or histidine labeling, etc. |
| Minute TM Non-Denatured Protein Solubilization Reagent | WA-010 | ELISA, immunoprecipitation/Co-IP, enzymatic activity determination and other applications. |
| Minute TM Protein Solubilization Reagent for MS | WA-011 | Trypsin digestion and subsequent mass spectrometry analysis |

About Evaluation of Isolated PM Proteins

Many researchers use Western blotting to access the purity of isolated membrane proteins. Some commonly used "cytosolic markers" are not exclusively cytosolic. For example, actin (1), GAPDH (2) and tubulin (3) are mainly cytosolic but they are also associated with plasma membranes. It's not surprising to detect week signals of these marker proteins in PM preps in certain cell and tissue types. For more info please refer to following publications:

- 1. Gruenstein E., et al. (1975). Actin associated with membranes from 3T3 mouse fibroblast and Hela cells. Journal of cell Biology. 64:223-234.
- 2. Terrasse R., et al. (2012). Human and pneumococcal cell surface glyceraldehydes-3-phosphate dehydrogenase (GAPDH) proteins are both ligands of human C1q protein. J. Biol. Chem. 287:42620-42633.



3. Wolff J. (2009). Plasma membrane tubulin. Biochemica et Biophysica Acta. (BBA)-Biomembranes 1788:1415-1433.

Troubleshooting

| Problem | Solution |
|--|---|
| Low protein yield | Increase starting cell numbers Increase incubation time to 10 min (step3) |
| ow protein activity Keep lysate cold/add protease inhibitors | |
| Retention of cell lysate in protein filter | Reduce amount of starting material or increase centrifugation |
| cartridge after 30 seconds of centrifugation | time to 2 min |
| Contamination of PM by cytosolic proteins | Wash PM pellet with 0.5 ml cold PBS containing 0.3 M |
| Contamination of FWI by cytosofic proteins | NaCl, pH 9.5 |