PRO-PREP™ Protein Extraction Solution

Cat. No.

17081

100 ml

DESCRIPTION

By using PRO-PREP $^{\text{TM}}$, proteins can be simply extracted from all kinds of cells an d tissues. The kit contains 5 kinds of protease inhibitors so it is possible to extra ct very highly purified proteins .

CONSIDERATION BEFORE USE

Usually detergent used in protein extraction consists of both hydrophobic tail and hydrophilic head as an amphiphilic molecule. The two parts are joined to form a micelle, that is, solubilized protein forming a lipid-detergent mixed micelle and tra nsmembrane protein forming a protein-lipid-detergent complex. The extent of mi celle formation is termed as CMC (critical micelle concentration), which is important for high efficiency as well as high purity of protein extraction. CMC is influenced by pH, temperature, ionic strength, multivalent ions of organic solvents, pur ity of detergent, and so on.

Depending on its ionic characteristics, a detergent can be categorized as ionic de tergent, non-ionic detergent, and Zwitterionic detergent. Ionic detergent can be fu rther classified into both cationic detergents: (SDS, LiDS and DOC) and anionic d etergent. Thus these are highly denaturant which have a specific property to isola te protein as a monomeric form and so often used in Western blot analysis and m easurement of molecular weight. Also non-ionic detergent such as Triton X-100 a re less protein denaturant and often employed in protein-protein interaction.

Zwitterionic detergent such as CHAPS have both negative and positive charge he ad at the same time, more effective in protein-protein interaction than non-ionic d etergent and its extent of protein denaturation is less than that of ionic detergent. It is very important to select the optimal buffer and detergent when extracting proteins

ADDED PROTEASE INHIBITORS

PMSF	- inhibits serine protease and thio protease - added in a working concentration of 1.0mM (174µg/ml)	
EDTA	- inhibits metaloprotease - added in a working concentration of 1.0mM	
Pepstatin A	- inhibits acid protease - added in a working concentration of 1μM (0.7μg/ml)	
Leupeptin	- inhibit serine protease - added in a working concentration of 1µM (0.5µg/ml)	
Aprotinin	- inhibit serine and thiol protease - added in a working concentration of 0.1μM (2.0μg/ml)	

CHARACTERISTIC

- When extracting proteins from cells or tissues, one doesn't necessarily apply a ppendix treatment.
- 2. Able to minimize protein extraction time within 20-30 minutes.
- 3. Protein stabilization buffer can make protein stable.
- 5. Extracted proteins are stable for more than 6 months when kept in -20~%.
- There is no absorbable error because there is no absorbable hindrances of PR O-PREP solution when measuring protein concentration.
- Very useful for protein separation in Western blot analysis because lonic deter gent turns protein into monomers.
- 8. Very useful for protein molecular weight analysis because denature protein int o a monomers
- Protein degradation is minimized by adding commonly used protease inhibitor and doesn't need to prepare protease inhibitor.

PROTOCOL (For Cells)

1. Preparation of cells.

Note: After preparation of adherent cell or suspension cell in 50ml tube, centr ifuge at 2,000-3,000rpm for 5 min. Then wash cells with PBS/DPBS (optional). After washing, count cells and use approximately 5x10⁶ cells. And then trans fer to the new 1.5ml tube.

2. Harvest the cell pellet by centrifuge at 13,000rpm for 10-20 seconds.

Note: After centrifugation, remove the remnant using a pipette.

3. Resuspend cells in 400µl PRO-PREP™ solution, and mix well.

Note: Depending on tissue types, one can vary volume of PRO-PREP™ solutio n. Generally add 400µl per 5x10⁶ cell, but determine the optimal amount of so lution according to cell size. Also, pipette carefully as the addition of PRO-PR EP™ solution can produce bubbles.

4. Induce cell lysis by incubation for 10-20 min on ice or freezer at -20 $^{\circ}$ C.

Note: PRO-PREP™ solution don't freeze at -20 °C, and it can be stabilize prot ein refraining protein degradation with protease inhibitor. Before incubating, it can also increase cell lysis using a syringe(optional). At this time, there appears bubbles, yet doesn't need to care because they disappear during centrifugation or incubation.

- 6. Measure of protein concentration.

Note: When measuring protein concentration by Bradford' method etc., PRO-P REPTM solution is made to have no absorbable hindrance, and so can decline a n absorbable error.

PROTOCOL (For Tissues)

1. Preparation of tissue about 10-20mg.

Note: After digging the interested tissue, transfer it to an appropriate tube. Ke ep the tissue fresh as much as possible.

2. Homogenize tissues in 600µl PRO-PREP™ solution.

Note: According to tissues, can be adding different addition of PRO- PREPTM s olution. Generally add 600µl per 10mg tissue, but determine to add the optim al amount of solution for each experiment. Also, when tissue is homogenized by homogenizer, there appears bubbles. If incubated or centrifuged, they will disappear. Doesn't need to care.

3. Induce cell lysis by incubation for 20-30 min on ice or freezer at -20 $^{\circ}\mathrm{C}$.

Note: PRO-PREPTM don't freeze at -20 °C, and it can be stabilized protein refr aining protein degradation with protease inhibitor. Before incubating, it can a lso increase cell lysis using a syringe(optional). At this time, there appears bu bbles, yet doesn't need to care because they disappear if centrifuged or incubated in freezer.

4. The following procedures are same as the PROTOCOL of cells.

TECHNICAL INFORMATION

Problem

In case of using tissue sample

EXPERIMENTAL INFORMATION

Yield

1. Protein extraction volume in various cells and tissues.

The conclusion of total protein volume used by PRO-MEASURE™ Protein Measure ment Solution after isolate protein used by PRO-PREP™ Protein Extraction Soluti on from various cells or tissues is average 2mg per 5 X 106 cells and 8-9mg per 1 Omg tissues.

1> Celll

Strain	Cell Number	Protein
K562 (human)	5 X 10 ⁶	2.04 mg
SNU 601 (human)	5 X 10 ⁶	1.89 mg
YAC 1 (mouse)	5 X 10 ⁶	2.10 mg
B16 (mouse)	5 X 10 ⁶	1.90 mg

2>Tissue

Strain	Cell Number	Protein
Spleen (mouse)	10 mg	9.01 mg
Kidney (mouse)	50 mg	9.91 mg
Lung (mouse)	10 mg	9.67 mg
Liver (mouse)	10 mg	6.59 mg

2. SDS-PAGE gel Electrophoresis conclusion

Total proteins were isolated from various cancer cell lines and tissues by the PR O-PREP ™ Protein Extraction Solution. We can extract the subcutaneous fat cont ained large amount of fat

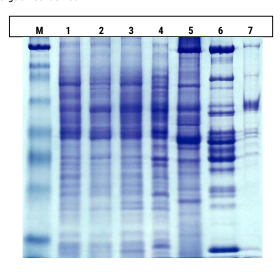


Fig. 1. SDS-PAGE gel Electrophoresis conclusion

This is the dying 15% SDS-PAGE gel electrophoresis conclusion used by coomassie bril liant dye after total protein extraction using by PRO-PREP™ Protein Extraction Solution i n various cancer cell line and tissue.

Lane M, PRO-STAIN (I) Marker; lane 1, K562; lane 2, SNU 1; lane 3, SNU 5; lane 4, B16; lane 5, rat Heart; lane 6, cow muscle; lane 7, Cow subcutaneous fat

TROUBLESHOOTING GUIDE

Recommendation

- First put tissue into a medicine pestle bowl and grind. The important thing is keep the

Possible Cause

	Cell number	medicine pestle bowl in refriger temperature.
Tiny of extraction protein. Regratable lysis.	Lysis hour	- Check properity of cell vol. and a following PRO-PREP™ Solution vol.
		Check did a right lysis method. Extend lysis hour and do vortexing
		every regular space during extract protein for higher lysis efficience.
Protein degradation		- Do not keep too long time in a room temperature while doing lysis with PRO- PREP™ Solution.
Too much bubble		 Check the extracted protein's keeping condition by use PRO-PREP™Solution and this product. Keep everything in 20 ℃.
comming out during pipetting		- Do not need worry because during doing
Freezed PRO-PREP Solution		incubation or doing centrifuge for lysis it can be disappear.
	In case of too low of keeping temperature	 Check a deep-freezer temp. If keeping temp, is lower than 20 °C, it can be freeze, at this point do not have problem if we use after melt it. However if we do again and again, it can have any problem, so keep a storing temp. as possible.
Occur clarity pellet in bellow part of PRO- PREP™ Solution		If we keep deep-freezer, PMSF among protease inhibitor should be clarity crystal condition. In case of this happen, after melt
	Protease Inhibitor	solution at room temperature in a minute, put ice solution for use cool temp. condition.
Con do ID 2		 If we cannot keep it at -20 °C, keep it separatively for need will be better for keep product's quality.
Can do IP ?		- Can do IP.

RELATED PRODUCTS

Product Name	Cat.No.
WEST-one™ Western Blot Detection System	16031 ~16033
WEST-ZOL™(plus) Western Blot Detection System	16021
PRO-MEASURE™Protein Measurement Solution	21011
SMART™ BCA Protein Assay Kit	21071
SMART™ Micro BCA Protein Assay Kit	21072
PRO-STAIN™Prestained Protein Marker (I)	24051
SMART™ Bacterial Protein Extraction Solution	17511