

PERK or GCN2 IP Western

Solutions

Lysis buffer 500ml

1% Triton x 100 5ml
150mM NaCl 15ml 5M
20mM Hepes pH 7.5 10ml 1M
10% glycerol 50ml
1mM EDTA 1ml 0.5M

*freshly add to lysis buffer just before use:

10mM tetrasodium pyrophosphate
100mM NaF
17.5 mM -glycophosphate
1mM PMSF
4 ug/ml Aprotinin
2ug/ml Pepstatin A

Ripa buffer (optional more stringent wash for IP)

10mM Tris pH 7.5
100mM NaCl (up to 500mM for more stringent wash)
1mM EDTA
0.5% Na-deoxycholate
0.1% SDS
1% Triton x100

2X laemlli

100mM Tris pH 6.8
20% glycerol
4% SDS
0.2% bromophenol blue
200mM DTT

50X Methanol free-transfer buffer 1L 50X:

1.9M glycine 144.1g
250mM Tris base 30.3g

- dilute to 1X for methanol free transfer

1X Methanol-transfer buffer 1L:

100ml 50X "methanol-free" transfer buffer (this is not a mistake, if methanol is added the tris-glycine buffer must be used at 5 times the strength as buffer with no methanol)
200ml methanol
700ml dH2O

Sample treatment and harvest

For Cells:

- 1) Treat cells eg. Thapsingargin 0.1 μ m 15 - 30 min.
Tunicamycin 0.5 - 2.5 μ g/ml, 3hrs
Amino acid deprivation (dMEM –leucine, 10% dialized FCS, Pen-strep,
glutamine)
- 2) Wash cells 2x ice cold PBS
1x ice cold PBS – EDTA (1mM)
- 4) Lyse cells in 500 μ l lysis buffer + phosphatase + protease inhibitors.
- 5) Scrape to 1.5ml tube, 5 min ice.
- 6) Clear extract by centrifugation at 14,000rpm 10 min, 4°.

For Tissues:

- 1) Harvest fresh tissue as quickly as possible.
- 2) Homogenize in 4ml/g tissue weight of lysis buffer using a motorized teflon and glass dounce.
- 3) When the tissue is completely homogenized, transfer to 1.5ml tubes and clear by centrifugation at 14,000rpm at 4C for 30 min.
- 4) Repeat centrifugation of supernatant for the best results. (keep repeating until extract is completely clear.)
- 5) Preclear extract with non-specific antibody by incubating for 1hr with 1 μ l nonspecific (or preimmune) antisera and 20 μ l washed protein A beads for 1hr at 4C with end-over-end rotation. Centrifuge at 14,000rpm and discard beads for 10 minutes. Use supernatant for IP.

Immunoprecipitation for both cells and tissues:

- 1) For each sample
Prebind 1 μ l Ab
15 μ l Protein A beads
50-100 μ l lysis buffer + phosphatase + protease inhibitors
15min-1hr RT on end-over-end rotator
- 2) Preclear extract with non-specific antibody by incubating with 1 μ l nonspecific (or preimmune) antisera and 15 μ l washed protein A beads for 1hr at 4C with end-over-end rotation. Centrifuge at 3000rpm and discard beads. Use supernatant for IP.
- 3) 1-2x wash Ab bound beads in lysis buffer or Ripa, resuspend in 50 – 100 μ l/sample aliquot to 1.5ml tubes.
- 4) Add tissue or cell extract to tubes with washed antibody bound beads.
- 5) End-over-end rotation @ 4°, 3 hours-Overnight.
- 6) Wash 3x 1ml lysis buffer + protease/phosphatase inhibitors. (or RIPA)
- 7) Wash 1x 1ml with PBS.
- 8) Remove all solution with 27g needle + syringe.
- 9) Add 15-20 μ l 1x laemmli buffer boil and load on gel (6 –7 % SDS PAGE).

Western

- 1) Transfer in methanol free transfer buffer diluted to 1X. (for PERK either + or - methanol is okay)
*****For phospho-GCN2 use methanol transfer buffer.**
- 2) Block 30 min in PBS, 5% Non Fat Dry (NFD)-milk (no tween).
*****For Phospho-GCN2 or Phospho-Perk specific westerns use 5% BSA (ICN cat #160069) in place of NFD milk in block and antibody incubations. BSA solution can be sterile filtered and reused to reduce costs.**
- 3) Primary antibody: in PBS- 5% NFD milk (or 5%BSA for phospho-specific antibodies), no tween,. (see below for times and temp)
- 4) Wash 3X PBS -0.02% tween 20.
- 5) Secondary antibody 1/3000 proteinA-HRP (best) or 1/5000 anti-rabbit-HRP 1hr RT. Dilute antibody in PBS- 5% NFD milk (or 5%BSA for phospho-specific westerns), no tween,.
- 6) Wash 3X PBS-0.02% tween 20.
- 7) ECL development.

Antibody dilutions:

PERK: 1/3000-1/5000

Incubate 1-2 hrs RT or 4C ON.

GCN2: 1/1000

Incubate 1-2 hrs RT or 4C ON.

P-GCN2: 1/500-1/1000

Incubate 4C ON.