

# In-gel digestion of proteins separated by SDS-PAGE

(Lamond Lab, January 2009)

- ☉ Perform all the pipetting steps in a laminar flow hood. We routinely do our digestions in our TC room hoods. Remember to wear gloves at all times, and don't scratch your head or any other part of your body during the procedure (if you do, change your gloves!) Keratin contamination is less of any issue *after* peptide extraction, as whole keratin proteins will not interfere with MS analysis.
- ☉ Peptides tend to stick to plastic surfaces on storage. We use 1.5 ml Eppendorf LoBind tubes (cat no. 2243108-1) that are designed for storing peptides at low concentrations.
- ☉ When preparing stock solutions, clean spatulas thoroughly with water and ethanol. Even better, simply tip the solid chemical into clean Eppendorf tubes.
- ☉ Acetonitrile (CH<sub>3</sub>CN) is light sensitive, so store in a dark bottle or tube wrapped in foil.

## Stage 1: If you are doing SILAC IP, first elute your IP samples from beads

**NOTE: To improve elution of proteins from beads and to save time during the digestion, we now elute in SDS and then reduce and alkylate the proteins prior to running them on a gel.**

1. *Elute proteins from beads (sepharose, agarose, dynabeads, etc.)* Add 1 volume of 1% SDS to the beads (e.g. 50  $\mu$ l of 1% SDS to 50  $\mu$ l of beads) and boil for 10 min at 95°C. Add 4 volumes of dH<sub>2</sub>O (e.g. 200  $\mu$ l of dH<sub>2</sub>O to 50  $\mu$ l of beads) and vortex well to elute proteins. Pellet the beads and collect the supernatant. This results in a more efficient release of proteins from the beads. **Note: Save the beads and add sample buffer directly (e.g. 20  $\mu$ l dH<sub>2</sub>O plus 20  $\mu$ l of 4X sample buffer) to elute any remaining proteins (can then run them on the same gel to see how much did not elute).**
2. *Speedvac the supernatant* back down to the original volume (e.g. 50  $\mu$ l), which will take the SDS concentration back to 1%. 45 min at 45°C is a good approximation.

## Stage 2: Reduction and Alkylation

- ☉ Reduction and alkylation of cysteine residues using DTT and IAA, respectively, improves the recovery of cysteine-containing peptides from in-gel digests and minimizes the appearance of unknown masses in MS analysis due to disulfide bond formation and side chain modification.
  - ☉ For complex samples (like cytoplasmic, nuclear or nucleolar extracts), we first add 4X NuPAGE LDS sample buffer from Invitrogen (NP0007) to the samples and then DTT and IAA as described below, avoiding step 3.
1. *Reduce the sample.* Add DTT to a final concentration of 10 mM (we use BDH 443553B) and boil for 1–2 min.  
(e.g. for 50  $\mu$ l, add 0.5  $\mu$ l of 1M DTT stock).
  2. *Alkylate the sample.* Add iodoacetamide (Sigma I1149) to a final concentration of 50 mM and incubate at room temperature in the dark for 30 min.  
(e.g. for 50  $\mu$ l, add 2.5  $\mu$ l of 1M IAA stock).
  3. Add 4X NuPAGE LDS sample buffer (a few  $\mu$ l is fine).

## Stage 3: Separating proteins on gels and excising bands

1. *Separate your protein sample by 1D PAGE.* We routinely use Novex precast gels. For a complicated sample we use 4–12% gradient gels, run them all the way (200V for 50 min) and cut the gel into 12–16 slices. For less complicated samples (e.g. IPs), we use straight percentage gels (usually 10% or 12%), run them halfway down (200V for 25 min) and cut the gel into 5–6 slices. *For IPs, we usually run the eluates in multiple lanes due to the large volume and for each slice cut across two or more lanes. This means the amount of gel in each tube is at least doubled and more buffer will need to be added to cover gel slices (especially important for the trypsin digestion step).*
2. *Stain the gel with Coomassie blue.* We routinely use the SimplyBlue SafeStain solution from Invitrogen (LC6060; protocol on the bottle). To minimize contaminants, do all staining steps in a sterile 14-cm tissue culture dish. Destain the gel thoroughly in dH<sub>2</sub>O (overnight is fine). The gel can be silver-stained at this stage.
3. *Scan the gel* before cutting out the bands. To do that, put the gel into the cover of the 14-cm dish and scan it. Print out the scanned image so that you can mark on it where you cut the bands. The gel can be returned to the dH<sub>2</sub>O-filled dish until ready to excise the bands.
4. *Excising the bands from the gel (4-5h for 96 bands).* For this step, we transfer the gel to a clean 14-cm tissue culture dish and cut away the unnecessary parts (top, bottom, MW marker lanes) with a sterile scalpel (e.g. Swann-Morton disposable scalpels, cat. no. 0511), leaving only the lanes in which you are interested. If you want to identify proteins in a single Coomassie-blue-stained band, excise the gel as close to the band as possible, with no excess around the band (to ensure that proteins you identify are from that one band).
5. *Mincing the gel bands.* Once you have the slices cut out for a particular sample lane (and marked on the printout of the scanned gel), cut each slice into 1 x 1 mm pieces using a fresh scalpel to cut and transfer each slice into a 1.5 ml Lobind Eppendorf tube. Try to work as quickly as possible, because the gel becomes stickier as it dries out. **The bands can be stored for later use at -20°C.** *Note: If your gel is silver stained, add 50 µl/band of 15 mM potassium ferricyanide/50 mM sodium thiosulphate (Farmers reagent - made fresh from 2X stock solutions) for 5–10 min until the band pieces go clear (i.e. until all the silver is removed).*

## **Stage 4: Destaining the gel bands**

**(about 3 hours for 32 tubes for Stage 4)**

1. *Wash the band pieces* with 300 µl of dH<sub>2</sub>O water for 15 min. Add 300 µl of CH<sub>3</sub>CN (Acetonitrile; Sigma A3396) and wash for a further 15 min.
2. *Remove the supernatant* (Use a P1000 tip with a P10 tip on the end, it is necessary because your gel pieces may be lost through the blue tips).
3. *Wash the band pieces* with 300 µl of 20 mM NH<sub>4</sub>HCO<sub>3</sub> (Sigma A6141, 1M aliquots at -20°C that are to be diluted in MS water) for 15 min. Discard the supernatant.
4. *Wash the band pieces* with 300 µl of 20 mM NH<sub>4</sub>HCO<sub>3</sub> / CH<sub>3</sub>CN (50:50 v/v) for 15 min. The gel pieces should shrink and look opaque. Discard the supernatant.
5. *If the band pieces are still blue, repeat the NH<sub>4</sub>HCO<sub>3</sub> and NH<sub>4</sub>HCO<sub>3</sub> / CH<sub>3</sub>CN washes.* (On adding NH<sub>4</sub>HCO<sub>3</sub>, the gel pieces should be restored to the original sizes and look transparent again.) You may crush the band pieces with a Teflon stick, but we found this step not essential in most cases, and it may increase keratin contamination.

6. Add 100  $\mu\text{l}$  of  $\text{CH}_3\text{CN}$  to dehydrate the band pieces for 5 min. The gel pieces should shrink and look completely white. Discard the supernatant.
7. Dry the band pieces in a Speedvac for 5 min.

## Stage 5: Digestion of band pieces

☉ Trypsin is a serine protease that specifically cleaves peptide bonds on the carboxyl side of lysine and arginine residues. However, cleavage can be blocked or slowed by proximal acidic, aromatic or proline residues, proline having the most significant effect. Peptide fragments with one missed cut are common and should be taken into consideration during mass analysis.

1. Add 50  $\mu\text{l}$ /band of 12.5  $\mu\text{g}/\text{ml}$  of modified trypsin in 20 mM  $\text{NH}_4\text{HCO}_3$ . We use Trypsin Gold from Promega (V5280), which is supplied as 100  $\mu\text{g}$  of powder. We resuspend it at 0.5  $\mu\text{g}/\text{ul}$  as our stock solution (i.e. add 200  $\mu\text{l}$  of 50mM acetic acid (see note) to the 100  $\mu\text{g}$  powder in the vial, and store it at  $-20^\circ\text{C}$  in 20  $\mu\text{l}$  aliquots). Trypsin is not stable especially when pure and in non-acidic condition, so we dilute down from this stock solution just before use. We routinely use 50  $\mu\text{l}$ /band of a freshly prepared 12.5  $\mu\text{g}/\text{ml}$  stock in 20 mM  $\text{NH}_4\text{HCO}_3$ .  
(for 1.8 ml of trypsin digestion buffer, add 45  $\mu\text{l}$  of Trypsin Gold stock to 1.755 ml of 20 mM  $\text{NH}_4\text{HCO}_3$ ).

**Note:** Preparation of 50mM acetic acid: We use MS grade acetic acid (100%). The molecular weight of acetic acid is 60.05 g / mol, and the density is 1.051 g / ml. Hence 57.13 ml = 60.05g = 1 mol, so 57.13 ml in 1 litre is 1M. To prepare a 50 mM solution, add 71.4  $\mu\text{l}$  in 25 ml of MS-water.

2. Allow bands to rehydrate in trypsin digestion buffer for 30 min. The gel pieces should be restored to the original sizes, and there should be JUST enough trypsin solution to cover all the gel pieces. If required, add more 20 mM  $\text{NH}_4\text{HCO}_3$  (minus the trypsin; more buffer will need to be added in the case of IPs since usually multiple lanes have been combined in one tube) to cover the band pieces. Keep a record of how much liquid you add per band, because you will need to know this for the first step of the extraction protocol (i.e. when you add an equal volume of  $\text{CH}_3\text{CN}$  to start extracting the peptides).
3. Incubate at  $30^\circ\text{C}$  overnight (>16h; no shaking).

## Stage 6: Extraction of peptides

Perform all the gel washing extraction steps on a shaking platform to ensure complete extraction of peptides.

1. Add an equal volume (e.g. 20 or 50  $\mu\text{l}$ ) of  $\text{CH}_3\text{CN}$  to the digest.
2. Incubate at  $30^\circ\text{C}$  for 30 min.
3. Transfer supernatant to a new clean Eppendorf LoBind tube. This supernatant contains the peptides you are going to analyze.
4. Add 50  $\mu\text{l}$  (or enough volume to cover the gel pieces) of 1% formic acid to the gel pieces. Prepare the 1% formic acid solution fresh in the fume hood, by adding 0.5 ml of 100% formic acid (BDH cat. no. 101155F) to 49.5 ml of  $\text{dH}_2\text{O}$ . Incubate for 20 min.
5. Transfer supernatant to the tube at step 3.
6. Repeat steps 4 and 5 once more.

7. Add 150  $\mu\text{l}$  of  $\text{CH}_3\text{CN}$  to the gel pieces. The gel pieces should shrink and turn white. Incubate for 10 min.
8. Transfer supernatant to the tube at step 3.
9. Speedvac to dry the peptides in the tube at step 3 completely (60°C is fine).
10. Resuspend the (invisible) pellet with 1% formic acid. To optimize solubilization of peptides, first add 10  $\mu\text{l}$  of 5% formic acid, vortex, and then add 40  $\mu\text{l}$  of  $\text{dH}_2\text{O}$ . Peptides should now be stored in the freezer.
11. For injection on the MS, thaw the peptides, spin hard to pellet any crap (e.g. 13,000g for 10 min) and transfer 15–20  $\mu\text{l}$  to a crimp vial (Kinesis ref. 20413) with a lid. Store the rest in the freezer.